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EXHIBIT A

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EXHIBIT A

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6	1.0 INTRODUCTION
7	
8	This document is a Remedial Action Plan (hereafter, "Remedial
9	Action Plan", "RAP", "Exhibit A" or the "Document") and describes
10	the work to be performed by the Settling Defendants at the
11	Environmental Conservation and Chemical Corporation ("ECC")
12	Superfund site as required by the attached Consent Decree
13	("Consent Decree" or "Decree"). This document is attached as
14	Exhibit A to, and is incorporated by reference into and made an
15	enforceable part of, that Decree.
16	
7	The purpose of this Exhibit A is to set forth those remedial
18	activities to be performed at the ECC site. The Settling
19	Defendants under the Consent Decree ("Settling Defendants") shall
20	arrange to have the work required hereunder performed by a
21	Contractor or Contractors ("Contractor") in accordance with the
22	requirements and specifications set forth herein.
23	
24	The components of the RAP as presented herein are compatible
25	with the proposed remedy for the adjacent Northside Sanitary
26	Landfill (NSL) site. As the remedial design is finalized for the
27	NSL site, the respective RAPs for ECC and NSL will be reviewed to
28	ensure compatibility of design and construction schedules for
29	each system. If any inconsistencies are identified, the Settling
30	Defendants shall consult with those performing the remedy at NSL,
31	and with EPA and the State to attempt to resolve any such
3 2	inconsistencies.

34 35 2.0 REMEDIAL ACTION PLAN 36 2.1 Elements of the RAP 37 38 2.1.1 Soil Vapor Extraction, Concentration and 39 40 Destruction 41 42 The objective of the soil vapor extraction activity is to remove 43 and destroy VOCs and selected base neutral/acid organics from the soils (as provided herein). 44 45 By systematically and uniformly moving air through the zone of 46 contamination, volatilization and hence removal of organics are . 47 accelerated. For the ECC site, air movement through the soil 48 will be controlled by a network of vertical trenches installed 49 50 throughout the zone of contamination. The process also involves the continuous extraction of organics-laden air from the trench 51 system and treatment of the air by activated carbon to remove the 53 organics. The organics so collected will then be destroyed offsite in conformance with applicable Federal and State 54 requirements. 55 56 The effectiveness of vapor extraction for organics removal from 57 the ECC soils was demonstrated during a pilot test conducted by 58 Terra Vac, an environmental consulting firm, in June, 1988. 59 description of the pilot test, including the results obtained, 60 was previously submitted to USEPA and the State of Indiana. 61 test showed an initial high organics extraction rate of 1.9 62 pounds per day per foot of trench that decreased over the course 63 of the pilot test to a steady state rate of approximately 0.25 64 pounds per day per foot of trench. Although the Terra Vac pilot

study provides the foundation for the system designed herein for 66 ECC, during the conceptual and preliminary engineering phase, 67 several engineering and operational enhancements were developed 68 which should improve overall performance and effectiveness of the 69 vacuum extraction system to be implemented under this Remedial Action Plan. These system enhancements are the result of 71 72 consultations among the following environmental consulting firms: ERM-North Central, Inc., Midwest Water Resource, Inc. (MWRI), and 73 Terra Vac, Inc. A summary of the key improvements and the associated measures employed for this enhanced vapor extraction 75 76 system are as follows: 77 78 Reduction of surface water infiltration 79 within the zone of treatment by construction 80 of the Resource Conservation and Recovery Act (RCRA) - compliant (Subtitle C) cover system; 81 82 Reduction in the volume of air required for 83 0 effective remediation by reducing air 84 infiltration into the vapor extraction system 85 by constructing the RCRA-compliant (Subtitle 86 87 C) cover; 88 89 Reduction of atmospheric discharges of 0 treated extraction air by reinjecting the air 90 through a network of injection trenches 91 installed as part of the vapor extraction 92 system; 93 94

Positive control (collection and removal) of

subsurface* till water encountered in the

zone of treatment by providing sufficient

95

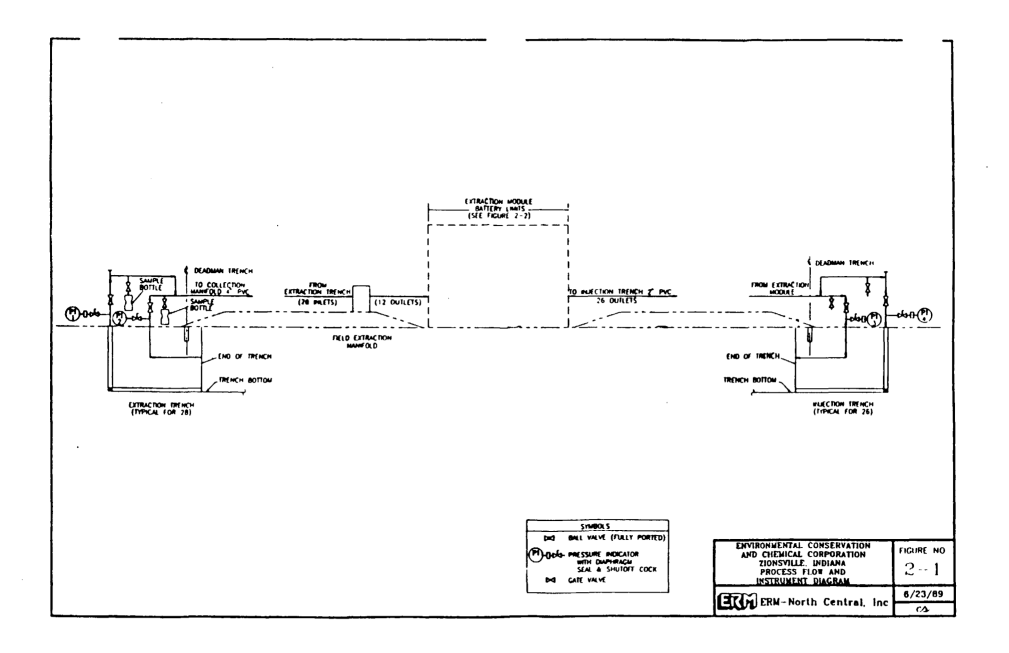
vacuum and/or supplemental air to remove 98 99 water which accumulates in the extraction trenches; and 100 101 Essentially uniform horizontal movement of 102 air through the zone of treatment resulting 103 104 in enhanced contact between the air and the VOCs in the soil during operation of the soil 105 vapor extraction system by utilizing a 106 network of injection and extraction trenches 107 in conjunction with the impervious cover 108 provided by the RCRA-compliant (Subtitle C) 109 cover system. 110 111 * For purposes of this document, "subsurface" water shall mean 112 "ground water", as defined at 40 CFR 260.10. 113 114 115 The following discussion and drawings show concepts and details 116 of the design and operation of the soil vapor extraction system. 117 The soil vapor extraction process is illustrated in Figures 2-1 118 and 2-2. The basic operation consists of extraction of air using 119 120 a single vacuum pump from a network of 28 extraction trenches 121 located throughout the site. Free liquid entrained in the air is 122 removed by gravity in an entrainment separator. Periodically, water which accumulates in the entrainment separator is pumped to 123 an on-site storage tank for subsequent transport to an off-site 124 facility for treatment as necessary, in accordance with 125 applicable Federal, State and local regulations. 126 From the vacuum

pump, air passes through the carbon adsorption system, which

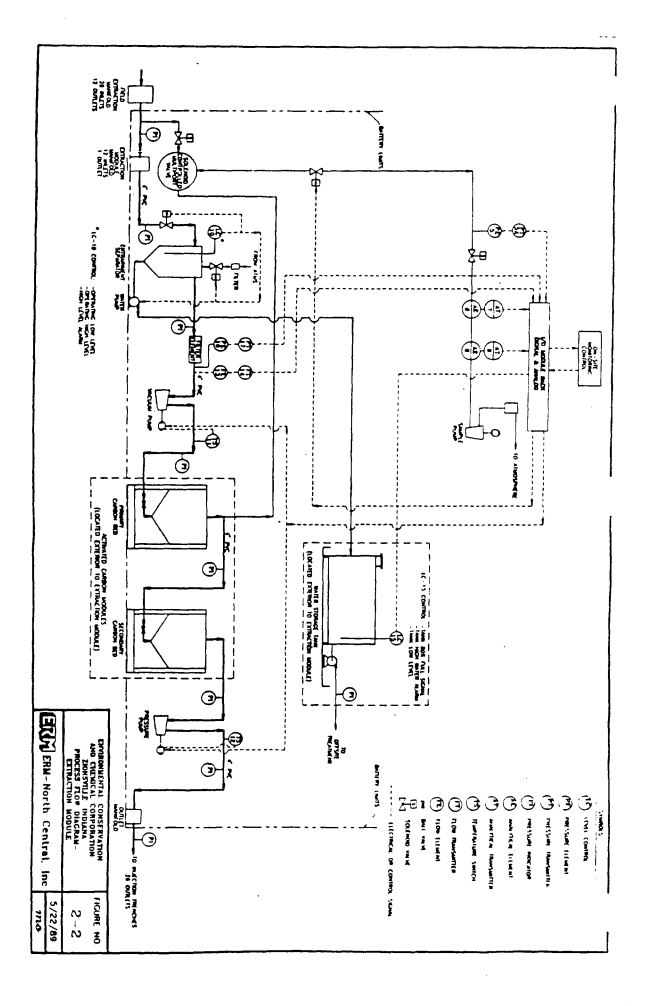
consists of two upflow carbon columns connected in series. Offgases from the carbon adsorption system are withdrawn by a pump

127

128

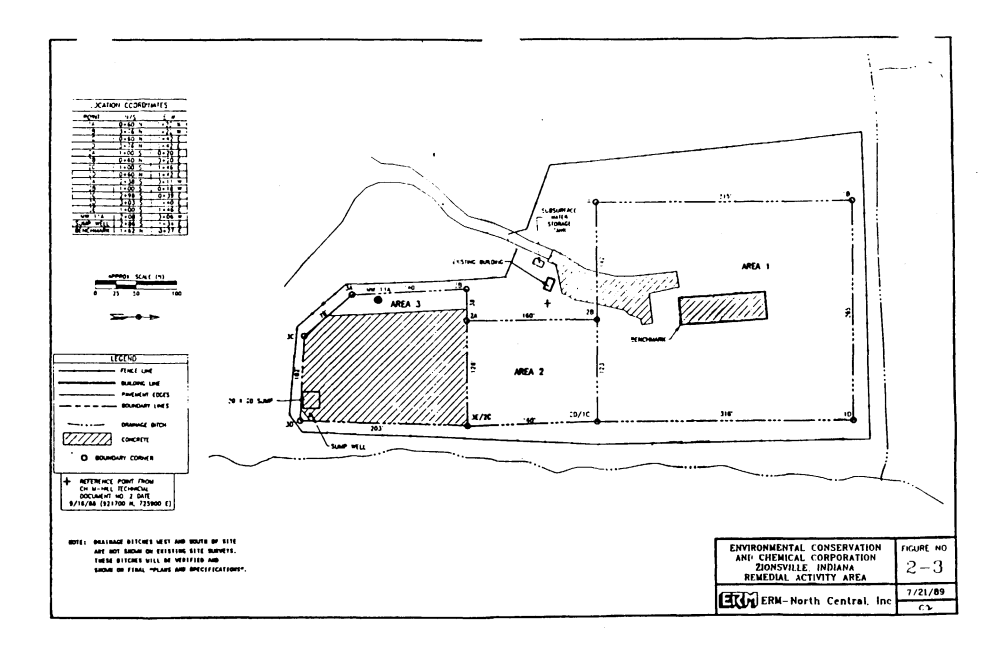


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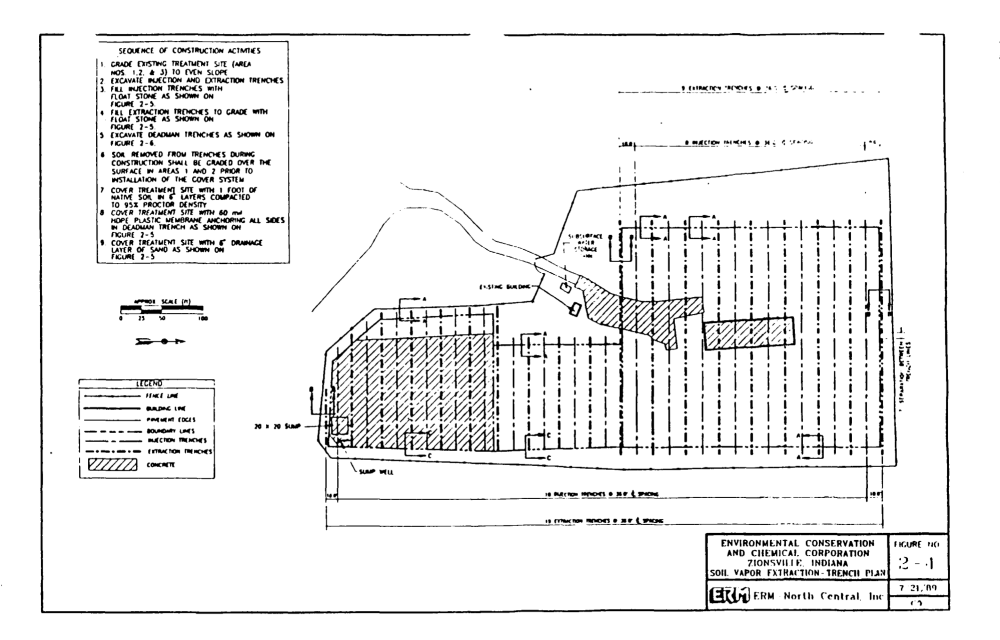
•

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which boosts the pressure and reinjects air into a network of 26
130
     injection trenches located throughout the site. Each injection
131
     trench is located between and parallel to a pair of extraction
132
    trenches. The injected air then migrates from the injection
133
     trench through the soil towards the extraction trench. As the
134
     air migrates through the soil towards the extraction trench, the
135
     organics are vaporized into the air stream. As described in
136
137
     Section 2.1.2, the RCRA-compliant (Subtitle C) cover will be
138
     placed over the entire trench network to prevent air and water
139
     infiltration into the system during operation.
140
141
     The major system components are:
142
               Extraction and injection trenches;
143
144
145
               Soil vapor extraction system;
          0
146
147
               Water collection system;
          0
148
149
               Carbon adsorption system;
          0
150
151
               Air injection system; and
152
153
               RCRA-compliant (Subtitle C) cover.
154
155
     A description of the design and operational features of each of
156
    these components is presented below.
157
158
          Extraction and Injection Trenches
159
    The area where remedial activity will occur is depicted in Figure
160
     2-3. The west boundary of Area 1 encompasses the area of ECC
```

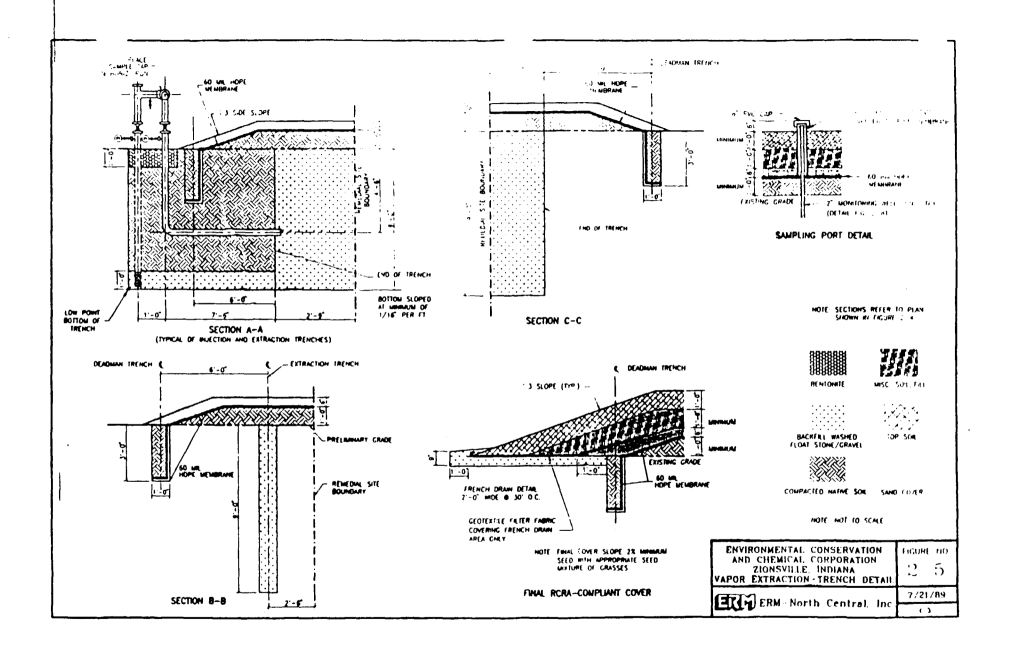


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activities that resulted in hazardous substances being released,
     as verified by an examination of aerial photographs, and
163
     coincides with a pre-existing earthen berm which formed the
164
     western boundary of ECC's water containment system for this area.
165
166
     The layout and construction details for the network of 28
167
168
     extraction trenches and 26 injection trenches are presented in
     Figures 2-4 and 2-5. Trench spacing will be 18 feet, and trench
169
     length varies depending on the configuration of the site.
170
171
     Construction details of extraction trenches and injection
     trenches are identical. By implementing minor above-ground
172
     piping changes, injection trenches can and will be utilized as
173
174
     extraction trenches. The work required under this Remedial
     Action Plan will initially involve using the original extraction
176
     trenches for extraction; at some point in the process, the
177
     extraction trenches will be converted to injection trenches, and
178
     vice versa, to ensure complete vapor extraction of the soil.
179
180
    All trenches are to be a minimum of 9-feet deep as measured from
     existing grade, and will be backfilled with washed "float" stone.
181
182
     The trench width will be 12-15 inches. The bottom elevation for
183
    both injection and extraction trenches will be sloped at a
184
    minimum of 1/16-inch per foot to a low point located at the water
     collection pipe as noted in Section A-A of Figure 2-5.
185
186
187
    Soil removed from the trench excavation will be spread over the
    surface of the facility prior to construction of the cover system
188
    and covered in accordance with the final RCRA-compliant (Subtitle
189
190
    C) cover detail illustrated in Figure 2-5. Soil removed from the
    trenches constructed in the areas of the concrete pad (Area 3)
191
102 will be spread over the surface in Areas 1 and 2 with trench
```

spoils from those areas.



194 195 Each trench will be equipped at one end with a vapor extraction pipe and a water collection pipe as illustrated in Section A-A of 196 Figure 2-5. Both pipes will be 4-inch diameter, Schedule 40 PVC. 197 Each pipe segment will be equipped with pressure/vacuum 198 indicator, isolating valve and sample tap. A "T" at the top of 199 the water collection pipe will permit the future installation of 200 201 air piping to air lift water from the trench network, if necessary. Individual 4-inch, Schedule 40 PVC pipes will be 202 routed from each extraction trench to the extraction module. 203 204 extraction module will be located adjacent to the existing concrete pad near the site entrance. Alternatively, two or three 205 206 extraction trenches will be manifolded together and conveyed to ! the extraction module via a 4-inch, Schedule 40 PVC pipe. 07 Injection trench piping is identical to the extraction trench 208 piping and, as previously described, will permit it to be 209 utilized as an extraction trench during the operation of the 210 211 vapor extraction system. To minimize field piping from the extraction module to the injection trenches, 4 to 8 injection 212 trenches will be manifolded together. Four-inch, Schedule 40 PVC 213 214 pipe will be used to convey air returned from the extraction 215 module to the injection trench. 216 The Sump Well installed by EPA will be backfilled with the 217 material used to backfill trenches (i.e, float stone) and a 4-218 inch PVC pipe will be installed between the Sump Well and the 219 220 nearest extraction trench, thereby tying the Sump Well directly into the vapor extraction system. The existing 20 ft. x 20 ft. 221 sump will be handled similarly, and will be dewatered prior to 222 223 installing the RCRA-compliant (Subtitle C) cover system. All water removed from this sump will be handled in accordance with ~24 applicable Federal, State and local requirements. .25



226 Soil Vapor Extraction System 227 228 229 The vacuum pump will have a nominal capacity of 500 standard 230 cubic feet per minute (SCFM) and will be capable of developing a 231 vacuum of 18 inches Hg. The normal operating vacuum is 232 anticipated to be 12 inches Hg. Based on MWRI's experience with soils characteristic of the ECC site and on the Terra Vac pilot 233 study results at the ECC site, the zone of influence at the 234 235 operating vacuum will be at least 40 feet (20 feet either side of 236 the trench). The pilot test results showed an initial radius of 237 influence of 15 feet during trench development. Under continuous operation, the radius of influence increased to about 20 feet. 238 The enhanced operating efficiency obtained by installing an impervious cover and injecting air will increase the radius of 40 influence to over 20 feet. To be conservative, a spacing between 241 trenches of 18 feet was selected. 242 243 The vacuum will be applied at the trench outlet and will be 244 245 distributed throughout the entire length and vertical dimension 246 of the trench. The highly porous backfill material used will 247 assure this uniform distribution of vacuum throughout the extraction trench. The reinjection pressure of air in each adjacent injection trench will be approximately 37.4 inches Hg 249 250 (1.25 atm). Therefore, the pressure differential and driving force for air movement between injection and extraction trenches under normal operating conditions is approximately 19.4 inches Hg 252

253 254 (0.65 atm).

255 The selection of the design air volume of 500 SCFM is based upon 256 MWRI's experience and is consistent with the Terre Vac pilot plant test results. The criteria established is to provide at

258 least one air volume change per soil pore volume per day. Based upon an area of treatment of 150,000 square feet, a depth of 259 contamination of 9 feet, and a soil porosity of 10%, 500 SCFM 260 261 exceeds the MWRI criteria by 400%. 262 263 The vapor extraction process will operate continuously and will 264 shut down automatically only in the event of an operating problem 265 or malfunction. The following are conditions which will shut down normal operating sequence of the vapor extraction system: 266 267 268 0 High vapor temperatures above the estimated acceptable range of 150 to 180°F prior to 269 activated carbon treatment; 270 1 Low vapor temperatures below the estimated , 2 0 acceptable range of 75 to 85°F prior to 273 274 activated carbon treatment indicating relative humidity above the estimated 275 276 acceptable range; 277 278 High water level in water entrainment 0 separator indicating operating problems with 279 280 liquid transfer operation; 281 High water level in subsurface water storage 282 0 283 tank; 284 High or low pressure conditions on vacuum or 285 0 286 injection pumps under normal operating 287 conditions; and 228

Power interruptions for the site.

During normal operation, vapor extraction will be stopped to
facilitate carbon vessel change out as described later in this
section and during transfer of water from the entrainment
separator to the on-site subsurface water storage tank, or to
conduct restart spike tests.

The air extracted from the system will be continuously monitored 297 298 by in-line instrumentation as shown on the process flow diagram (Figure 2-2) and described on Table 2-1 (Instrument Summary 299 300 Sheet). The capability will exist to sample individual trench 301 exhausts or the combined air stream. Sample taps will be provided to collect vapor samples for detailed chemical analysis. 302 0.3 The on-line instrumentation will consist of a photoionization detector (PID) and moisture analyzer. The vacuum pump, controls 304 305 and instrumentation will be located in the Vapor Extraction Module Building. 306

307

Water Collection System

308 309

310 The high vacuum vapor extraction system selected will be capable 311 of entrainment and movement of water which accumulates in the 312 extraction trenches. Any free liquid in the extracted vapor will be separated by gravity in an entrainment separator located in 313 314 the Vapor Extraction Module Building. A level control system will be utilized to control the removal of water which 315 accumulates in the entrainment separator as required. 316 317 separator tank is equipped with a vacuum breaker system which 318 will open the tank to the atmosphere to permit water to be 319 transferred by pump from the separator to an on-site water 30 storage tank as necessary. The time required to make this transfer will depend upon the equipment supplied by the vapor 21

TABLE 2-1 ERM-NORTH CENTRAL, INC. INSTRUMENT SOMMARY SHEET

CLIENT: Environmental Conservation and Chemical Corporation

PROJECT NO.: 9041

and the second s

DATE:	2/28/89 REVISED: 5/23/89	Trenes	
TAG NO.	SERVICE	MOUNTING PANEL FIELD	SPEC FLOW SHEET DIA.
PI	Pressure Indicator	x	2-1 2-2
PI-1	Pressure Indicator	((
thru	with diaphram	· } x	. 2-1
PI-4	Seal and shutoff cock		
PE-5	Pressure sensing element		
PT-5A	Pressure transmitter	x	2-2
AE-6	Moisture sensing element	x	2-2
AT- 7	Moisture transmitter	X	2-2
AE-8	Volatile organics detector and quantifier	x	2-2
AT-9	Volatile organics quantified signal transmitter	x	2-2
LC-10	3-point water level control and alarm	x	2-2
TS-11	Gas temperature sensor with high level system shutdown switch	x	2-2
TS-12	Gas temperature sensor with high level system shutdown switch	x	2-2
FE-13	Gas flow measuring element	x	2-2
FT-14	Gas flow signal transmitter	X	2-2
LC-15	3-point water level control and alarm	x	2-2
PE-16	Pressure sensing element	x	2-2
PI-17 .	Pressure transmitter	x	2-2

322 extraction system vendor selected.

323

324 The size of the storage tank will be sufficient to store the 325 liquids, considering the off-site handling/treatment option 326 selected. If water collected from the soil vapor extraction system is to be discharged to the Northside Sanitary Landfill 327 328 (NSL) pipeline, a 1,000-gallon storage tank will be used; or if water collected is to be hauled off-site by tank truck for 329 330 disposal, a 10,000-gallon tank will be used. The tank will be equipped with level measurement and control to advise operating 331 personnel to the status of liquid accumulation in the storage 332 tank. Periodically, the contents of the water storage tank will ! 333 need to be removed. The removed water will either be sent to the 334 Indianapolis POTW via the NSL pipeline or truck, or to another off-site facility for handling and treatment as necessary, in 336 accordance with applicable Federal, State and local regulations. 337

338

339 340

Carbon Adsorption System

341

From the water entrainment tank, the air passes through a 342 343 particulate filter preceding the vacuum pump. The pressure drop 344 across the filter will be monitored and used as the signal for determining servicing of the filter element. The exhaust from 345 the vacuum pump will be piped directly to a two-stage carbon 346 347 adsorption system (primary and secondary). This system will consist of two vessels in series each containing approximately 348 349 1,800 pounds of granular activated carbon. The organics 350 contained in the extracted air will be adsorbed on the activated The moisture content of the air stream will be less 351 than 50% relative humidity and temperatures will be approximately 150°F, both acceptable for efficient operation of carbon

354 adsorption. 355 356 During the initial phases of operation, when organics 357 concentrations in the air stream will be highest, the carbon 358 capacity for the organics is expected to be about 25% by weight. 359 During the latter phases of remediation as organic concentration 360 of vapor decreases, the projected carbon capacity for organics 361 will range between 10-15% by weight. Based upon an assumed total 362 mass of organics of about 5,000 pounds (Appendix A), the total quantity of activated carbon required for the entire remediation 363 364 program is 25,000 pounds. This equates to fourteen 1800-pound 365 carbon vessels for the entire program. The actual amount of carbon used will depend upon the total mass of organics extracted 366 67 during operation of the soil vapor extraction system and the 368 carbon adsorption capacity. 369 370 The vapor from the primary carbon vessel will be monitored 371 frequently (approximately once per hour) by an on-line PID 372 analyzer. When the PID analyzer detects organic vapor in the air stream between the primary and secondary carbon vessels, the 373 374 vacuum extraction system will shut down automatically to permit 375 the removal and replacement of the "spent" primary carbon vessel. 376 An operator will be alerted to this condition, and will 377 disconnect the primary carbon bed from service. The spent carbon 378 vessel will be removed and replaced by a carbon vessel containing 379 fresh activated carbon. The unit previously serving as the 380 secondary carbon bed will become the primary carbon bed and the 381 unit just placed in operation will be the secondary carbon bed. 382 Once this switch is complete, the soil vapor extraction system 383 (i.e., vacuum pump and injection pump) will be restarted, and the

system operation resumed. The arrangement of two activated carbon vessels in series (i.e., primary and secondary) will

permit optimal utilization of the activated carbon, and efficient 386 387 capture of the organics.

388

The spent carbon vessels will be stored on-site. The vessels 389 390 will be stored on the existing concrete pad adjacent to the vapor 391 extraction module building, inside the fenced area. approximate location of this area is shown in Figure 2-4. 392 393 inlet and outlet connections to each vessel will be capped and 394 sealed appropriately. Periodically when a truckload quantity of vessels has accumulated, and at the conclusion of the vacuum 395 extraction program, the vessels containing the spent carbon will 396 397 be transported in accordance with applicable Federal, State and 398 local requirements to an off-site facility where the carbon will 199 be regenerated by high temperature incineration, and in the process, the organics adsorbed on the carbon will be destroyed. 00

401

Air Injection System

402 403

404 The exhaust air from the secondary carbon bed will be piped to 405 the injection pump located in the extraction module building. 406 The injection pump will be capable of delivering 500 SCFM at 10 407 psig (1.65 atm). The discharge from the injection pump will be 408 distributed to the 26 injection trenches via a system of 409 manifolds. Control of the injection pump will be interlocked 410 with the vacuum extraction pump. The pipe at each injection 411 trench will be equipped with a pressure/vacuum gauge so that injection pressure at the trench can be periodically monitored. 412 413

- 414 During the soil vapor extraction program, the injection trenches will be utilized as extraction trenches and vice versa. 415
- 416 be accomplished by minor above ground manifold piping
 - modifications. It is also planned that as the Cleanup Standards 7

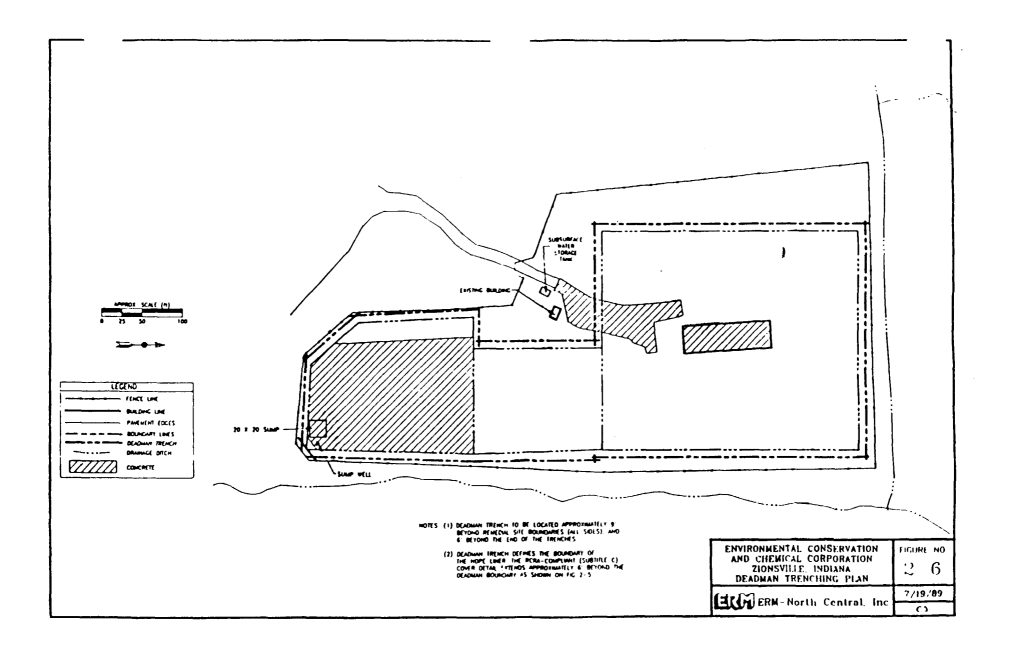
set forth in Table 3-1 below are met for individual trench 418 "areas", the corresponding extraction and injection trenches will 419 be isolated from the extraction and injection operation by 420 closing the shut off valves located at each trench. 421 422 permit the soil vapor extraction system to concentrate on any remaining areas which have not fully achieved the Cleanup 423 Standards specified in Table 3-1, thereby accelerating cleanup of 424 425 those areas. 426 RCRA-Compliant (Subtitle C) Cover 427 428 The operation of the vapor extraction system will be enhanced by 429 the installation of the RCRA-compliant (Subtitle C) cover over 430 the entire site. Details and a schedule for installation of the 31 final RCRA-compliant (Subtitle C) cover are presented in Section 432 2.1.2. 433 434 435 Miscellaneous 436 437 0 Each extraction trench is equipped with two sample taps, one on the vacuum pipe and one 438 439 on the water collection pipe. Each of these 440 taps can be fitted with a sample bottle for the collection of free moisture. 441 442 Electrical service required for the site 443 0 444 remediation work is anticipated to be 3phase 460 volt. Total electrical demand will 445 446 be approximately 100 KVA. Power distribution will be to the extraction module building. 447 Operating voltage for the extraction and 148 injection pumps is anticipated to be 460 49

450		voics. A 110 voic supply will be provided
451		for miscellaneous site lighting, equipment,
452		instrumentation and controls. Power
453		distribution to any site construction and
454		office trailers will also be provided.
455		
456	•	Prior to construction of the trenches, the
457		following activities will be conducted:
458		
459		1. The existing buildings within the
460		area currently fenced will be
461		demolished and properly disposed of
462		off-site;
463		
464		2. The existing tanks removed and
465		properly disposed of off-site; and
466		
467		3. The site will be graded to fill
468		existing depressions and to
469		eliminate any sharp grade changes.
470		
471		2.1.2 RCRA-Compliant (Subtitle C) Cover
472		
473	The RCRA-	compliant (Subtitle C) cover illustrated in Figure 2-5
474	will cons	ist of a minimum of 1-foot of compacted, highly
475	impermeab	le native soil, a continuous welded 60 millimeter high
476	density po	olyethylene (HDPE) plastic membrane, a minimum 6-inch
477	layer of o	compacted sand for drainage, 1 to 3 feet of
478	miscellane	eous soil/fill material and 1 foot of top soil to
479	support ve	egetation. The final grading plan will ensure a minimur
180	slope of 2	2%. The native soil used will be the silty clay till
81	available	in the area, which can and will be compacted by

standard methods to 95% proctor density. If soil from the 482 neighboring NSL Facility borrow area is not available, material 483 with similar performance will be obtained by Settling Defendants 484 485 from another source. 486 487 To provide a perimeter seal of the HDPE membrane, a 1-foot wide, 3-foot deep "deadman trench" will be installed around the site 488 boundary (Figure 2-6). The HDPE membrane will be draped into 489 this trench. The trench will then be backfilled and compacted 490 with native soil (silty clay till) to 95% proctor density. The 491 cover will extend approximately 6 feet beyond the deadman trench 492 as noted on Figure 2-6 and detailed on Figure 2-5. 493 494 195 As previously described, the material excavated from the trenches 496 will be graded uniformly throughout trench areas 1 and 2 and incorporated into the top layer of existing surface soil prior to 497 the construction of the RCRA-compliant (Subtitle C) cover as 498 499 shown in Figure 2-5. 500 501 The RCRA-compliant (Subtitle C) cover will be installed over the entire site, including the concrete pad. Prior to operation of 502 503 the soil vapor extraction system, the following components of the RCRA-compliant (Subtitle C) cover will be installed: (1) 1-foot 504 505 minimum compacted native soil; (2) a 60 mil HDPE membrane; and 506 (3) 6 inches of sand. Prior to installation of the remaining 507 components of the cap, Settling Defendants shall ensure that the 508 aforesaid components of the cap meet the aforesaid specifications. The remaining components (1-foot minimum 509 miscellaneous soil/fill, 1-foot minimum topsoil and appropriate 510 vegetation) will then be installed in accordance with the 511 512 schedule presented in Section 5.0. At completion of the soil

vapor extraction program all surface piping will be removed from

13ر



```
that time the extraction and injection trench piping may be cut
515
516
     off at the current grade, filled with grout, and covered with a
     minimum of 1 foot of topsoil, which will be vegetated.
517
     Vegetation which will be established shall include fibrous,
518
519
     shallow, laterally growing roots, such as grass (which may
520
     include red fescue and Kentucky blue grass).
521
     The Settling Defendants shall conduct periodic inspections and
522
523
     shall repair the cap as necessary to ensure its integrity in
     accordance with the time periods set forth in 40 CFR Sections
525
     265.117 and .118 or 329 I.A.C. Sections 3-21-8 and -9.
526
 27
     2.1.3 Access Restrictions
28د
529
     Access restrictions to be implemented by the Settling Defendants
530 will consist of a fence around the site perimeter and the posting
531
     of warning signs. In addition, Settling Defendants will use
532
     "best efforts", as that term is used in Section X A. of the
533
     Decree, to have recorded appropriate restrictions with the County
534
    Recorder's Office prohibiting: (a) usage of the site for
535 excavation and development; (b) usage of ground water from the
536
     saturated till and the underlying sand and gravel; and (c)
537
     installation of new water wells other than monitoring wells.
538
539
540
     2.1.4 Subsurface and Surface Water Monitoring
541
542
    The monitoring activities will:
543
- 4
               Detect the presence of the VOCs, base
```

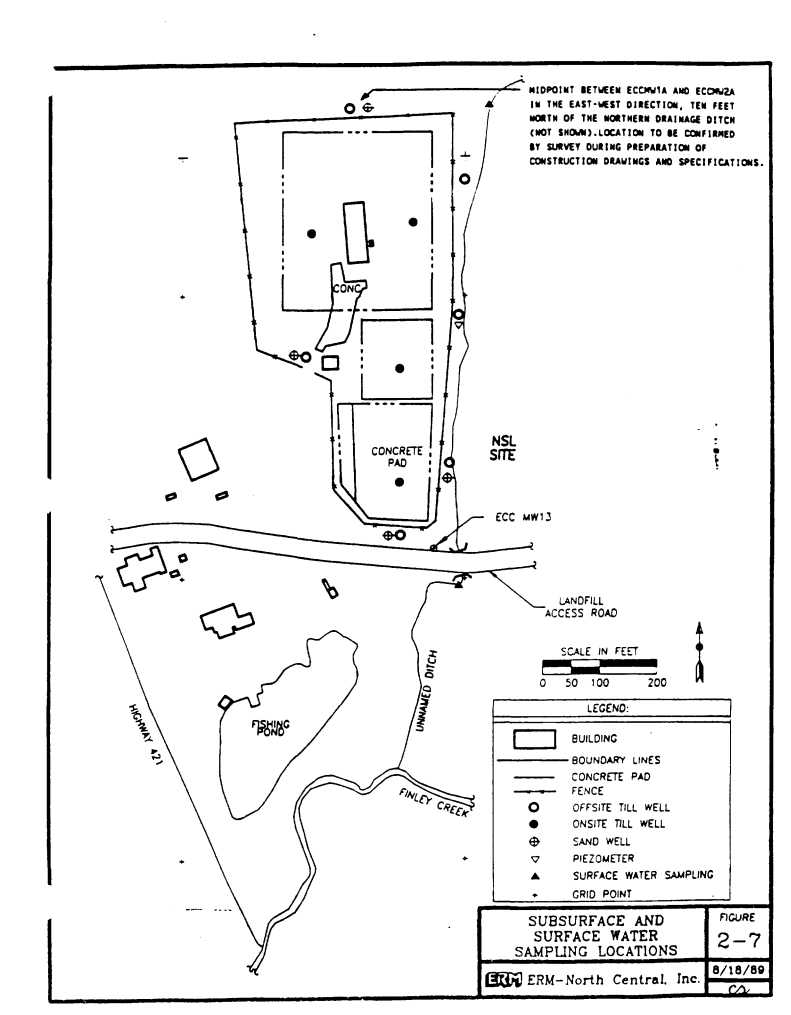
514 the site in addition to any equipment, buildings or trailers. At

neutral/acid organics, PCBs, and heavy metals

546 specified in Table 3-1 in the subsurface and 547 surface water during and after vapor extraction; and 548 549 550 Provide information to determine the 0 551 effectiveness of the soil vapor extraction 552 program. 553 554 Two types of subsurface water monitoring systems will be installed under this Remedial Action Plan. The first is an on-555 site till monitoring system consisting of four wells screened in 556 557 the saturated zone of the till. The location of these on-site ~58 till wells is shown in Figure 2-7. Sampling results from the onsite till wells will be compared to the Acceptable Subsurface 560 Water Concentrations in Table 3-1 or the Applicable Subsurface 561 Water Background Concentrations of Table 3-1 ("Applicable 562 Subsurface Water Background Concentrations"). 563 564 Samples from the on-site till monitoring wells will be collected 565 at the beginning of the soil vapor extraction operation and 566 quarterly thereafter until completion of the soil vapor 567 extraction program. Monitoring will be continued on a semiannual basis as specified in Section 4.0. Every time samples are 568 569 collected from the on-site wells, the soil vapor extraction system will be shut down to allow water, if any, to stabilize 570 within the till. Samples collected from the on-site wells will 571 572 be analyzed for those parameters listed under Acceptable Subsurface Water Concentrations in Table 3-1. 573 574 575

الغيار أأراف كالمواصر بمارات والرابوان المجتمع فللمعطف استنفاه فيراث والماماة

The second type of subsurface water monitoring system consists of off-site wells screened in the till and offsite wells screened in

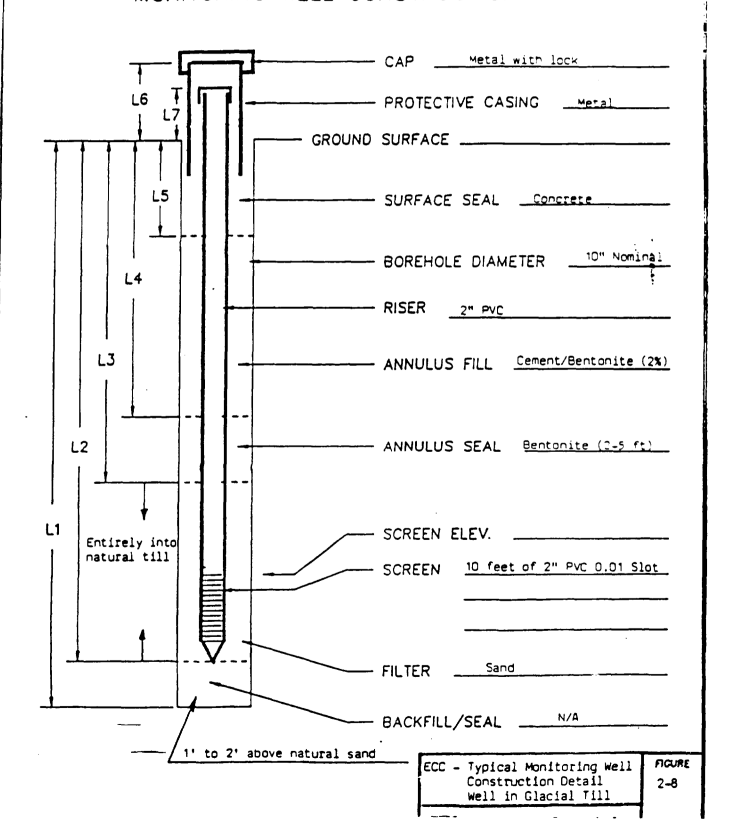


the sand and gravel. Sampling results from these wells will be 578 579 used to determine compliance with the Acceptable Stream Concentrations in Table 3-1 or the Applicable Surface Water 580 Background Concentrations of Table 3-1. 581 This second subsurface water monitoring network will consist of ten (10) new wells, 582 583 which will be located around the periphery of and downgradient from the ECC site, and one existing monitoring well, ECC MW-13 584 (Figure 2-7). In addition, a piezometer will be installed on the 585 586 east side of the site, as shown in Figure 2-7, to aid in defining 587 the direction of subsurface water flow in the sand and gravel. 588 Six (6) wells will be installed in the till, completed in the saturated zone, and four (4) wells will be completed in the sand 589 and gravel unit underlying the saturated surface till. 590

91

92د All wells (on-site and off-site) will be constructed of 2-inch PVC pipe. Screen length will vary for each well. 593 Total depth for the wells completed in the till will be 1-2 feet less than 594 595 total depth to the contact between the till and underlying sand and gravel. Wells completed in the sand and gravel will screen 596 597 the total thickness of that sand and gravel unit. Screens will 598 have a 0.01 inch opening. Wells will have a sand pack to one 599 foot above the top of screen and a bentonite grout to the ground surface. For the on-site till wells, a sampling port will be 600 fabricated in the HDPE membrane which will prevent infiltration 601 602 of air via these monitoring wells during operation of the soil vapor extraction system. A detail of this sampling port is shown 603 on Figure 2-5. Figures 2-8 and 2-9 illustrate well construction 604 605 details for the subsurface water monitoring wells in the till and in the sand and gravel, respectively. Details of the piezometer 606 607 construction are shown in Figure 2-10. The location of the 178 monitoring wells is based on the subsurface water elevation contours shown in Figure 2-11. 3

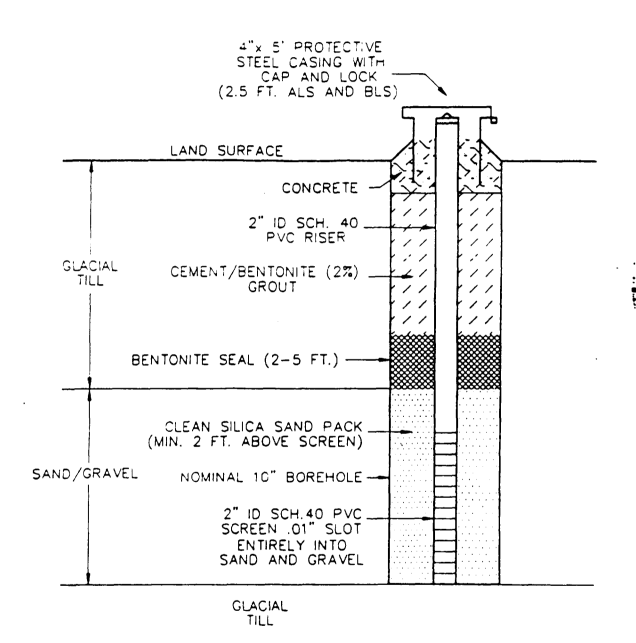
MONITORING WELL CONSTRUCTION



MONITORING WELL CONSTRUCTION — CAP <u>Metal with lock</u> - PROTECTIVE CASING _____Metal____ - GROUND SURFACE _____ **L5** — SURFACE SEAL ___Concrete_ 10" Nominal - BOREHOLE DIAMETER - RISER 2" PVC ANNULUS FILL Cement/Bentonite (2%) L3 L2 --- ANNULUS SEAL Bentonite (2-5 ft) In upper tills L1 Entirely into - SCREEN ELEV. natural sands 2" PVC 0.01 Slot SCREEN (Entire depth of sand and gravel Sand FILTER - BACKFILL/SEAL N/A ECC - Typical Monitoring Well FIGURE

Construction Detail well in Sand & Gravel

2-9



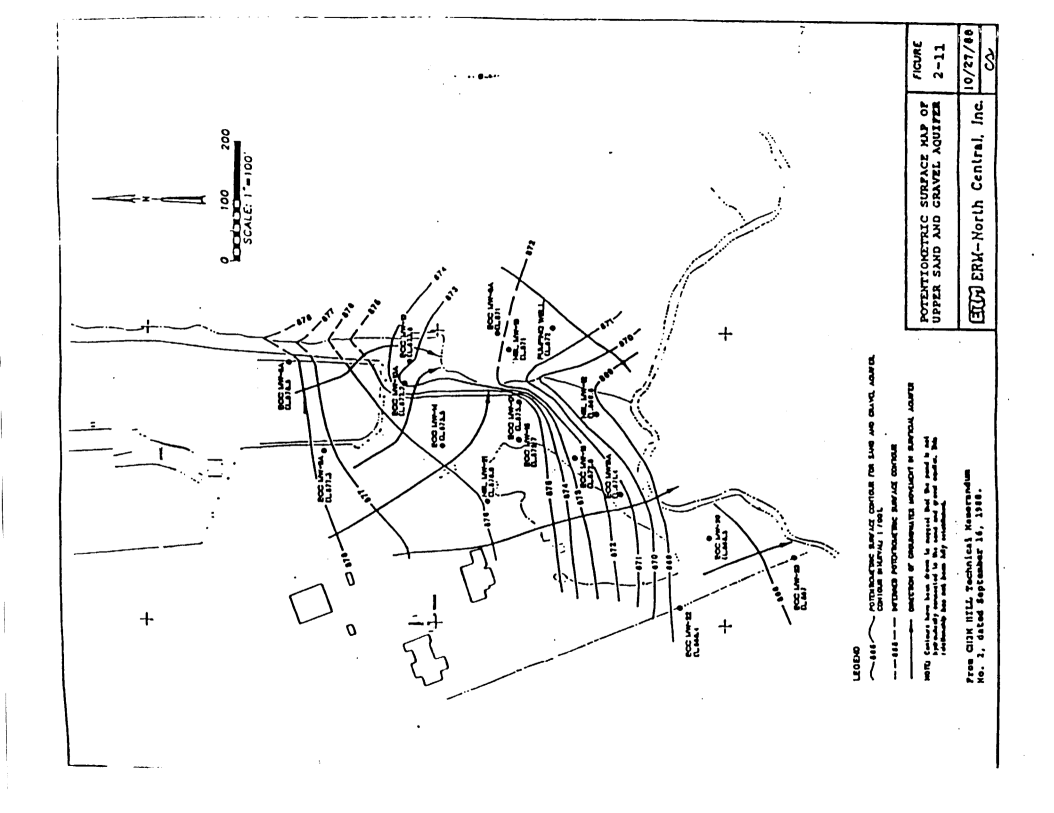
NOT TO SCALE

ECC-TYPICAL PIEZOMETER
CONSTRUCTION DETAIL
INSTALLED IN SAND AND GRAVEL

FIGURE 2-10

ERM-North Central, Inc.

8/18/89 *M*O



611 Samples from the off-site wells will be collected quarterly during operation of the vapor extraction system and analyzed for 612 613 the parameters with Acceptable Stream Concentrations in Table 3-614 Monitoring will be continued on a semi-annual basis as specified in Section 4.0. 615 616 617 The surface water will be monitored by sampling the Unnamed Ditch 618 just upgradient and just downgradient of the ECC site as depicted 619 in Figure 2-7. Surface water will be sampled at the same frequency as the off-site subsurface water and analyzed for the. 620 621 parameters with Acceptable Stream Concentrations in Table 3-1. 622 523 624 3.0 REMEDIAL ACTION CLEANUP STANDARDS 625 This section presents site-specific Cleanup Standards to be used 626 627 at the ECC site as the criteria for determining completion of 628 remedial action. The Cleanup Standards in this section are the 629 basis for establishing the criteria for Soil Cleanup 630 Verification presented in Section 4.2, and the Post-Soil Cleanup 631 Verification Compliance Monitoring in Section 4.3. If Soil Cleanup Verification as defined in Section 4.2 and the 632 633 subsections thereof is not achieved within 5 years of commencing 634 operation of the soil vapor extraction system, the Additional Work provisions of Section VII of the Consent Decree will apply. 635 636 637 638 639 640 3.1 Cleanup Standards 641

642	The follow	ring Cleanup Standards will be met for successful
643	completion	of the soil vapor extraction program:
644		
645	•	Acceptable Soil Concentrations shown in Table
646		3-1 will be achieved according to the
647		procedure discussed in Section 4.2.3 of
648		Exhibit A;
649		
650	•	Acceptable Stream Concentrations or
651		Applicable Surface Water Background
652		Concentrations shown in Table 3-1 will be
653		achieved in Unnamed Ditch south of and
654		adjacent to ECC;
555		
656	. 0	Acceptable Subsurface Water Concentrations
657		or Applicable Subsurface Water Background
658		Concentrations shown in Table 3-1) in the
659		on-site till wells will be achieved; and
660		
661	0	Acceptable Stream Concentrations or
662		Applicable Surface Water Background
663		Concentrations shown in Table 3-1 in the
664		off-site wells will be achieved.
665		
666	The term "	Table 3-1" wherever referred to or used in this Exhibit
667	A and in t	he Consent Decree includes the Footnotes on pages 2 and
668	3 of 3 of	that table.
669		
670		
671	3.2	Calculation of Cleanup Standards
672		

73 Table 3-1 sets forth the ECC site specific Cleanup Standards and

TABLE 3-1 (Page 1 of 2)
SITE-SPECIFIC ACCEPTABLE CONCENTRATIONS
ENVIRONMENTAL CONSERVATION AND CHEMICAL CORPORATION (ECC.) SITE

	1			
	Accepta			
	Subsurface		Acceptable Stream	Acceptable Soil
	Concentration		• • •	
Campounds	(ug/	-	(ug/l)	(ug/kg)
NO. LELLE COMMUNICATION COMMUNICATION		• • • • • • •	•••••	••••••
VOLATILE ORGANICS (VOCs):	7 500			.00
Acetone	3,500	-		490
Chlorobenzene		MCLGP		10,100
Chloroform	100		15.7	2,300
1,1-Dichloroethane	0.38			5.7
1,1-Dichloroethene		MCL	1.85	120
Ethylbenzene		MCLGP	3,280	234,000
Methylene Chloride	4.7		15.7	20
Methyl Ethyl Ketone		LDWHA		75
Methyl Isobutyl Ketone	1,750			8,900
Tetrachioroethene	0.69		8.85	130
Totuene	•	MCLGP	3,400	238,000
1,1,1-Trichloroethane		MCL	5,280	7,200
1,1,2-Trichloroethane	0.61		41.8	22
Trichloroethene	_	MCL	80.7	240
Total Xylenes	440	MCLGP		195,000
BASE NEUTRAL/ACID ORGANICS:				
Bis(2-ethylhexyl)phthalate	2.5		50,000	
Di-n-Butyl Phthalate	3,500		154,000	
Diethyl Phthalate	28,000		52,100	
Isophorone	8.5			
Naphthalene	14,000		620	
Phenol	1,400	RB	570	9,800
INORGANICS:				
Antimony	14	RB		
Arsenic		MCL	0.0175	
Barium	1,000			
Beryllium	175	RB		
Cadmium	10	MCL		
Chromium VI	50	MCL	11	
Lead	50		10	
Nanganese	7,000	RB		
Nickel	150	LDUHA	100	
Silver	50	MCL		
Tin	21,000	RE		
Vanadium	245	RB		
Zinc	7,000	RB	47	
Cyanide	154	LDWHA	5.2	
PESTICIDES/PCBs:				
PCBs	0.0045	RB (7)	0.000079 (7,8)	

TABLE 3-1 (Page 2 of 2) SITE-SPECIFIC ACCEPTABLE CONCENTRATIONS ENVIRONMENTAL CONSERVATION AND CHEMICAL CORPORATION (ECC SITE)

.

NOTES:

- (1) RB = Risk-based standard. U.S. EPA, Draft RCRA Facility Investigation Guidance, 1987.
 - MCL = Drinking water Maximum Contaminant Level. 40 CFR 141
 - MCLGP = Drinking water MCL goal, proposed. U. S. EPA Superfund Public Health Evaluation Manual, update of November 16, 1987.
 - LDWHA = Lifetime drinking water health advisory. U.S. EPA, Superfund Public Health Evaluation Manual, update of November 16, 1987.
- In the event that higher concentrations than those set forth for any parameter in this column are present in the upgradient subsurface water in the till and/or sand and gravel according to the procedure specified below, then those higher upgradient subsurface water concentrations and not the values set forth in this table shall constitute the Acceptable Subsurface Water Concentrations within the meaning of this Exhibit A and the Consent Decree. Those upgradient subsurface water concentrations are referred to in this Exhibit A as "Applicable Subsurface Water Background Concentrations." Twelve subsurface water samples will be taken from existing or new well locations, approved by EPA, over at least a 12 month period in areas upgradient of the site. The exact procedure, location of wells, and schedule for collecting and analyzing the samples will be approved by EPA, after consultation with the State, prior to its implementation. Subsurface samples for inorganics and PCB analysis will be filtered. For each parameter, the analytical results from the 12 samples will be analyzed using standard statistical procedures. The mean and standard deviation will be calculated, and all nondetects will be assigned a value equal to 1/2 the EPA-approved quantification limit. For purposes of this Document, "Applicable Subsurface Water Background Concentrations" is defined as two (2) standard deviations above the calculated mean of these 12 samples.
- (3) Stream Criteria, from Table 1 of the Record of Decision for the site, September 25, 1987.
- (4) In the event that higher concentrations than those set forth for any parameter in this column are present in the upstream surface water, then those higher upstream concentrations and not the values set forth in this table shall constitute the Acceptable Stream Concentrations within the meaning of this Exhibit A and the Consent Decree. Those higher upstream surface water concentrations are referred to in this Exhibit A as

"Applicable Surface Water Background Concentrations." Twelve surface water samples will be taken from Unnamed Ditch upstream of the site over at least a 12 month period. The exact procedure, location of samples, and schedule for collecting and analyzing the samples will be approved by EPA, after consultation with the State, prior to its implementation. For each parameter, the analytical results from the 12 samples will be analyzed using standard statistical procedures. The mean and standard deviation will be calculated, and all non-detects will be assigned a value equal to 1/2 the EPA-approved quantification limit. For purposes of this Document, "Applicable Surface Water Background Concentrations" is defined as two (2) standard deviations above the calculated mean of these 12 samples.

- (5) Acceptable Soil Concentration is based on ingestion of subsurface water at the site boundary, assuming a dilution of leachate to subsurface water of 1:196 (Appendix B).
- (6) The Acceptable Soil Concentrations, within the meaning of this Exhibit A and the Consent Decree, will be achieved when the arithmetic average of the 20 soil sample results for each parameter, assigning all non-detect results a value of one-half the detection limit, do not exceed the values set forth in this table by more than 25 percent.
- (7) So long as the EPA-approved quantification limit for PCBs in water is above the acceptable subsurface water and stream concentrations for PCBs, compliance with the Acceptable Subsurface and Stream Concentrations for PCBs will be determined as follows: all subsurface and surface water sample results for PCBs must be below the EPA-approved quantification limit for PCBs (at the time compliance is determined).
 - (8) Modified from Superfund Public Health Evaluation Manual, October, 1986, EPA 4/540/1-86/060, OSWER Directive 9285.4-1.

the procedure for determining Applicable Surface Water and 674 Subsurface Water Background Concentrations. The equations for 675 calculation of the risks, supporting data and complete references 676 677 are included in Appendix B. 678 679 The calculation of risk-based concentrations shown in Table 3-1 680 follows the procedures presented in the USEPA Draft RCRA Facility Investigation (RFI) Guidance, July, 1987, and in the USEPA 681 682 Memorandum on Interim Final Guidance for Soil Ingestion Rates, 683 January 27, 1989. In accordance with this latter reference, the soil ingestion rate for risk calculation was either 0.1 grams of 684 685 soil per day for a 70 kilogram person for 70 years (for compounds) with potency factors) or 0.2 grams of soil per day for a 17 686 · 7 kilogram child for 5 years (for compounds with reference doses). In accordance with the RFI Guidance document referenced above, **J**8 689 the ingestion rate used for the risk calculation was 2 liters of 690 water per day by a 70 kg person for 70 years. 691 Three columns of data, corresponding to Acceptable Concentrations 692 693 for Subsurface Water, Stream and Soil are presented in Table 3-1. Additionally, Applicable Subsurface Water Background 694 695 Concentrations, and Applicable Surface Water Background 696 Concentrations are defined in Table 3-1. The Acceptable Subsurface Water Concentrations are based on either drinking 697 water standards or criteria (Maximum Contaminant Level [MCL], 698 proposed Maximum Contaminant Level Goal [MCLGP], lifetime 699 drinking water health advisory [LDWHA]) or the appropriate risk-700 based concentration. These limits assume, as a worst case, that 701 the subsurface water in the till could be utilized as a lifetime 702 source of drinking water. However, the use of the subsurface 703 water in the till as a source of drinking water was rejected as unlikely in the ECC Remedial Investigation (RI), page 6-22.

706 a result, the use of drinking water standards and risk-based 707 standards based upon daily, long-term human consumption of the till water for Cleanup Standards under this Remedial Action Plan 708 709 represents an extremely conservative assumption when the real-710 life risks, if any, presented by the ECC site are considered. 711 712 The Acceptable Stream Concentrations are taken from the Record of 713 Decision (ROD) for the site, dated September 25, 1987. 714 715 The Acceptable Soil Concentrations in Table 3-1 are based on the 716 lowest of the risk-based concentrations for soil or subsurface water ingestion, from Tables B5 and B6. 717 718 719 Table 3-2 presents the compounds detected in soils at the site at 720 levels above the Acceptable Soil Concentrations specified in Table 3-1. Table 3-3 shows the vapor pressure and solubility of 721 722 these compounds. 723 724 3.3 Additional Work 725 726 If Additional Work is required under Section VII of the Consent Decree, Settling Defendants shall perform the following 727 728 additional work at the site unless the parties agree otherwise: 729 Maintain the RCRA-compliant (Subtitle C) 730 731 cover and the access restrictions. 732 733 Construct a subsurface water interception 0 734 trench around the south and east sides of the 735 ECC site as depicted in Figures 3-1 and 3-2. 736

Collect and transport subsurface water

37

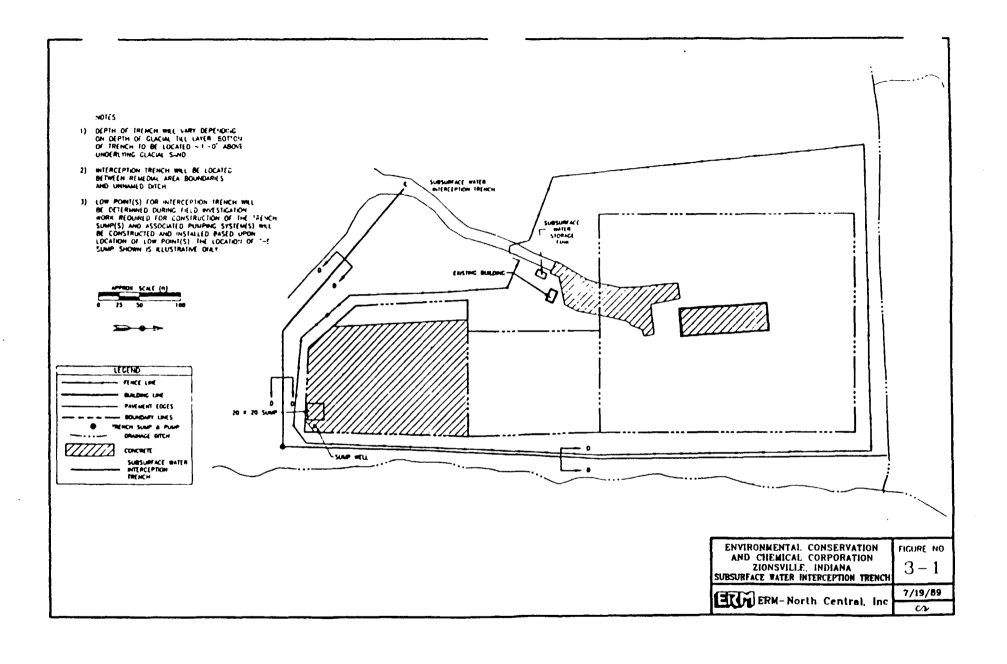


TABLE 3-2
COMPOUNDS DETECTED IN THE SOIL AT CONCENTRATIONS
ABOVE THE ACCEPTABLE SOIL CONCENTRATIONS (1)

Compound	Acceptable Soil Concentration (ug/kg)	Maximum Detected Concentration (ug/kg)
VOLATILE ORGANICS (VOCs):		
Acetone	490	650,000
Chloroform	2,300	2,900
1,1-Dichloroethane	5.7	35,000 ·
1,1-Dichloroethene	120	380 :
Ethylbenzene	234,000	1,500,000
Methylene Chloride	20	310,000
Methyl Ethyl Ketone	75	2,800,000
Methyl Isobutyl Ketone	8,900	190,000
Tetrachloroethene	130	650,000
Toluen e	238,000	2,000,000
1,1,1-Trichloroethane	7,200	1,100,000
1,1,2-Trichloroethane	22	550
Trichloroethene	240	4,800,000
Total Xylenes	195,000	6,800,000
BASE NEUTRAL/ACID ORGANICS:		
Phenol	9,800	570,000

⁽¹⁾ Acceptable Soil Concentrations are determined in accordance with Footnotes 5 and 6 of Table 3-1.

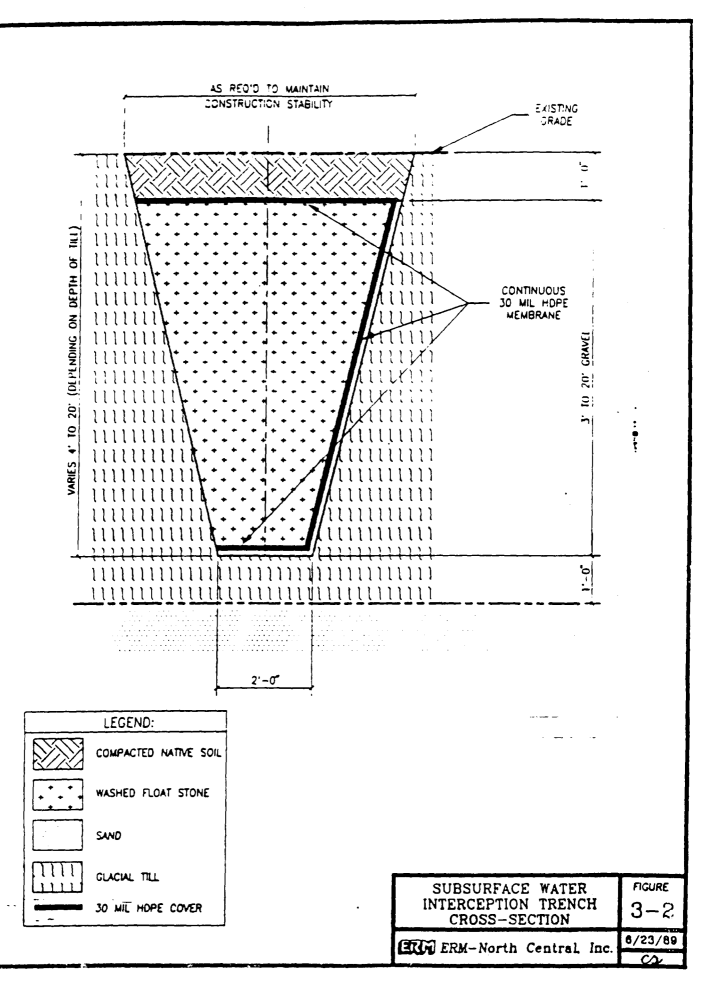
TABLE 3-3
CHEMICAL PROPERTIES OF ORGANIC COMPOUNDS
DETECTED IN THE SOILS AT CONCENTRATIONS
ABOVE THE ACCEPTABLE SOIL CONCENTRATIONS (1)

Compound	Solubility (ug/l)	Vapor Pressure (mm Hg)
VOLATILE ORGANICS (VOCs):		
Acetone	1,000,000,000	270
Chloroform	8,200,000	151
1,1-Dichloroethane	5,500,000	182
1,1-Dichloroethene	2,250,000	600
Ethylbenzene	152,000	7
Methylene Chloride	20,000,000	362
Methyl Ethyl Ketone	268,000,000	77.5 .
Methyl Isobutyl Ketone	17,000,000	. 6
Tetrachloroethene	200,000	17.8
Toluene	535,000	28.1
1,1,1-Trichloroethane	4,400,000	123
1,1,2-Trichloroethane	4,500,000	30
Trichloroethene	1,100,000	57.9
Total Xylenes	198,000	10
BASE NEUTRAL/ACID ORGANICS:		
Phenol	93,000,000	0.341

(1) Acceptable Soil Concentrations are determined in accordance with Footnotes 5 and 6 of Table 3-1.

REFERENCES:

- U.S. EPA, "Superfund Public Health Evaluation Manual," 1986.
- U.S. EPA, "Water-Related Environmental Fate of 129 Priority Pollutants," December 1979.



738 intercepted in this trench to the
739 Indianapolis POTW (via the NSL pipeline or
740 tank truck), or provide other appropriate
741 handling and treatment of such water in
742 accordance with applicable Federal, State and
743 local requirements.

2ر o Subsurface water will continue to be removed and handled in this manner until "confirmed" analytical results from two consecutive, semi-annual subsurface water samples collected from the interception trench show that the Acceptable Stream Concentrations in Table 3-1 or Applicable Surface Water Background Concentrations have been met, unless the Parties to the Decree otherwise agree.

o Semi-annual monitoring of off-site wells and surface water will continue for five years after the Acceptable Stream Concentrations in Table 3-1 or Applicable Surface Water Background Concentrations have been achieved.

If "confirmed" analytical results from two consecutive semi-annual samples collected during the 5 years of off-site monitoring in either the surface water or the wells indicate that the same parameter exceeds its Acceptable Stream Concentration or Applicable Surface Water Background Concentration at the same monitoring point, then subsurface water collection and treatment will be reinstituted.

770 As used in this section and in section 4.3 below, the term 771 772 "confirmed" shall permit the Parties to demonstrate that an 773 analytical result is not accurate as a result of errors in sampling, analysis, or evaluation or that it otherwise 774 mischaracterizes the concentration of a parameter. 775 procedures used to obtain "confirmed" data shall include 776 777 reanalysis, resampling and the analysis of only undiluted samples 778 if a concentration is qualified with a "J" (estimated 779 concentration). If after reanalysis and/or resampling using an 780 undiluted sample the concentration of a compound is still qualified with a "J", then the result produced from undiluted 781 782 samples will be used. "B" qualified samples results will be considered as "confirmed" data only if the concentrations in the 33 sample exceed ten times the maximum amount detected in any blank 84 785 for the media being analyzed.

786

787

789

788 4.0 REMEDIAL ACTION VERIFICATION AND COMPLIANCE MONITORING

790

The soil vapor extraction system described herein is designed to 791 achieve the cleanup standards for VOCs as presented in Table 3-1

792 and phenol. The time required to accomplish this removal depends

on the type of compound and soil, air flow rate and temperature, 793

794 and on an efficient diffusion of air through the soil pores.

795 time required for treatment was estimated using a vapor

796 extraction model, as described below and in Appendix C.

797 Monitoring of vapor from the combined vapor stream and from

798 individual trenches, as described below, will also be used to

799 estimate completion of the soil vapor extraction system

000 operation. Afterwards, verification of soil cleanup will be accomplished by: (1) soil vapor monitoring of restart spikes;

803 sampling ("Soil Cleanup Verification"). 804 805 Compliance monitoring will consist of sampling of surface water 806 in Unnamed Ditch, and sampling of subsurface water in off-site 807 till and sand and gravel monitoring wells and on-site till monitoring wells ("Compliance Monitoring"). 808 809 810 811 4.1 Estimation of Completion of Vapor Extraction 812 System Operation 813 A computer model which simulates the vapor extraction system was 814 15 used to estimate the time required for removal of the maximum detected soil concentrations to the Acceptable Soil 16 Concentrations specified in Table 3-1. Appendix C summarizes the 817 characteristics of the model and the data used. Based on the 818 819 model results, the Settling Defendants expect that after one 820 year of operation, all the VOCs and phenol will be below the Acceptable Soil Concentrations in Table 3-1 in a "worst case" 821 822 soil element which contains all the compounds at their maximum 823 detected concentrations. 824 825 The vapor extraction system is designed to permit vapor samples 826 to be obtained from each individual extraction trench and from 827 the combined vapor stream from all operating extraction trenches. 828 829 The combined vapor flow will be sampled daily during the first 830 week of operation, weekly for the following 4 weeks, and monthly thereafter. Samples will be analyzed for VOCs listed in Table 3-831 1 and phenol. Also, the vapor flow rate will be monitored and 832 recorded to provide sufficient data to calculate the mass of

(2) on-site subsurface till water monitoring; and (3) soil

802

organics removed from the soils and the effectiveness of the 834 system. These data will also aid in estimating the treatment 835 836 time remaining, based on the calculated mass extraction rate (lbs/day) of the VOCs listed in Table 3-1 and phenol. 837 838 839 Vapor samples from individual extraction trenches will be 840 collected at the beginning of the vapor extraction system operation to establish a baseline of organics removal per trench. 841 842 These samples will be analyzed for the VOCs listed in Table 3-1 843 and phenol. Once the mass rate extracted per day is reduced to 5 844 percent of the initial week's rate, additional vapor samples of individual trenches will be collected at least every two months, : 845 to determine when individual extraction trenches can be shut 846 The criterion for shutting down individual trenches will 47 be that two consecutive air samples from an individual trench 48 show vapor concentrations to be in equilibrium with the 849 Acceptable Soil Concentrations in Table 3-1. Table 4-1 shows the 850 851 soil vapor concentrations in equilibrium with the Acceptable Soil 852 Concentrations for the VOCs listed in Table 3-1 and phenol.

Appendix D presents the methodology used to arrive at these

854 855 856

853

4.2 Soil Cleanup Verification

equilibrium vapor concentrations.

857

Verification of soil cleanup will be established when each of the following is met: (1) the soil vapor from the restart spike tests shows compliance with the calculated soil vapor concentrations in equilibrium with Acceptable Soil Concentrations for the VOCs listed in Table 3-1 and phenol ("Soil Vapor Criterion"); (2) onsite till wells show compliance with the Acceptable Subsurface Water Concentrations specified in Table 3-1 or Applicable Subsurface Water Background Concentrations ("Onsite Till Water

TABLE 4-1
SOIL VAPOR CONCENTRATIONS IN EQUILIBRIUM WITH ACCEPTABLE SOIL CONCENTRATIONS (1)

	Soil Vapor Co	ncentration (3)
Compound (2)	(mg/l)	ppmv
VOLATILE ORGANICS (VOCs):		
Acetone	0.613	254
Chloroform	2.46	496
1,1-Dichloroethane	0.014	3.4
1,1-Dichloroethene	2.045	515
Ethylbenzene	37	9,316
Methylene Chloride	0.079	22.4
Methyl Ethyl Ketone	0.039	13
Methyl Isobutyl Ketone	0.685	233
Tetrachloroethene	0.116	16.8
Toluene	107	36,556
1,1,1-Trichloroethane	8.29	2,819
1,1,2-Trichloroethane	0.0060	1.1
Trichloroethene	0.39	71.5
	- · · - •	
Total Xylenes	26.2	4,794
BASE NEUTRAL/ACID ORGANICS:		
Phenol	0.0053	1.4

⁽¹⁾

Acceptable Soil Concentrations are determined in accordance with Footnotes 5 and 6 of Table 3-1. Compounds above acceptable soil concentrations in Table 3-1 to be removed by vapor extraction. From Appendix D. (2)

⁽³⁾

Criterion"); and (3) soil samples show compliance with the 866 Acceptable Soil Concentrations as specified in Table 3-1 ("Soil 867 Sample Criterion"). If Soil Cleanup Verification is not 868 established, vapor extraction will be restarted. If after five 869 years from the initial commencement of soil vapor extraction (or 870 871 sooner as permitted in the Decree), Soil Cleanup Verification has 872 not been established, then the Additional Work provisions of 873 Section VII of the Consent Decree will apply.

874

4.2.1 Soil Vapor Criterion

875 876

Once the combined vapor flow and individual trench vapor samples show concentrations of Table 3-1 VOCs and phenol at or below their respective equilibrium soil vapor concentrations shown in Table 4-1, the "restart spike" method on the combined vapor flow will be used to demonstrate that the Soil Vapor Criterion for Soil Cleanup Verification has been achieved.

883

884 The "restart spike" method consists of periodically shutting down 885 and restarting the vapor extraction system. By shutting down the system, equilibrium conditions between the vapor space within the 886 soil and any remaining organics amenable to vapor extraction 887 888 within the soil matrix are re-established. Therefore, when the 889 vapor extraction system is restarted, the initial organics 890 concentration in the extracted gas will be higher than under normal operation. 891

892

The restart spike procedure will include shutting down the vapor extraction system for a period of three days. Upon restarting the vapor extraction system, all extraction and injection trenches will be operated as during normal operation. A sample

37 of the combined soil vapor will be collected over a five-hour

period starting 30 minutes after restarting the vapor extraction system. This sample will be representative of the soil vapor concentrations in equilibrium with the soil concentrations, because at 500 SCFM, the vapor extraction system will exchange one pore volume of soil every five hours.

903

904 The Soil Vapor Criterion will be met when analyses of soil vapor samples collected from four consecutive restart spikes conducted once every two weeks show that concentrations of VOCs and phenol in Table 3-1 are at or below equilibrium soil vapor concentrations shown in Table 4-1 and therefore by calculation can be shown to be at or below the Acceptable Soil
910 Concentrations in Table 3-1.

911 912

4.2.2 On-site Till Water Criterion

913

914 Samples of the subsurface water from the on-site till monitoring 915 wells will be collected quarterly during operating of the soil 916 vapor extraction system. The most recent quarterly sampling 917 results from the four on-site till water monitoring wells 918 following demonstration that the Soil Vapor Criterion has been 919 achieved (Section 4.2.1) will be used to demonstrate that the On-920 site Till Water Criterion for Soil Cleanup Verification has been 921 achieved.

922

This criterion will be met when analyses of the water samples collected from each of the four on-site till wells show that the concentrations for parameters with Acceptable Subsurface Water Concentrations in Table 3-1 are at or below the Acceptable Subsurface Water Concentrations in Table 3-1 or Applicable Subsurface Water Background Concentrations.

:9

4.2.3 Soil Sample Criterion

930 931

Once the Soil Vapor Criterion and Onsite Till Water Criterion for 932 933 Soil Cleanup Verification have been demonstrated as defined 934 above, a total of twenty (20) soil samples from areas selected by EPA and the State will be collected. These twenty (20) will be 935 936 selected as follows: sixteen soil samples will be from "hot" 937 spot areas and four non-background samples will be from randomly 938 selected points elsewhere onsite. The total number of soil 939 samples used to demonstrate that the Soil Sample Criterion for 940 Soil Cleanup Verification will not exceed 20. Each soil sample will be analyzed for the VOCs in Table 3-1 and phenol. 941 942 Verification of this criterion for all VOCs in Table 3-1 and ~43 phenol relative to the Acceptable Soil Concentration in Table 3-If the results from this initial round of soil samples verify .4 945 that the Acceptable Soil Concentrations in Table 3-1 have been met, then the Soil Sample Criterion for Soil Cleanup Verification 946 947 will have been achieved.

948

949 In the event that the soil sampling results do not verify that 950 the Acceptable Soil Concentrations as defined in Table 3-1 have 951 been met, and the soil vapor extraction system is operated for an 952 additional period of time, an additional 20 soil samples must be taken in the same approximate locations (i.e., within a 3-foot 953 954 radius) as the initial sample locations. Results from this 955 second sampling will be analyzed using the identical procedure outlined above to verify that the Acceptable Soil Concentrations 956 in Table 3-1 as described in Footnote 6 of Table 3-1 have been 957 958 If the results from any subsequent round of soil samples demonstrate that the Acceptable Soil Concentrations in Table 3-1 959 have been met, then the Soil Sample Criterion for Soil Cleanup 960 Verification will have been achieved.

963 4.3 Post Soil Cleanup Compliance Monitoring 964 Once Soil Cleanup Verification has been achieved as prescribed in 965 Section 4.2, sampling of off-site till wells, on-site till wells, 966 off-site sand and gravel wells and surface water will be 967 968 conducted for seven years on a semi-annual basis. 969 970 Off-site wells and surface water will be analyzed for the parameters with Acceptable Stream Concentrations in Table 3-1. 971 Onsite wells will be analyzed for parameters with Acceptable 972 Subsurface Water Concentrations in Table 3-1. 973 974 975 If "confirmed" analytical results from two consecutive semiannual samples collected during the Compliance Monitoring period ·76 977 indicate that the same parameter exceeds its Cleanup Standard 978 (or the Applicable Surface Water or Subsurface Water Background 979 Concentration) at the same monitoring point, then the Additional 980 Work provisions of Section VII of the Decree will apply. conditions set forth in the preceding sentence do not occur, 981 monitoring will be discontinued at the end of the Compliance 982 Monitoring period and the provisions of Section XXVI of the 983 Decree will apply. 984 985 5.0 MISCELLANEOUS PROVISIONS AND SCHEDULING 986 987 The following documents have been submitted to EPA and the State 988 for review and approval by EPA: (1) Health and Safety Plan, (2) 989 Field Sampling Plan, and (3) Quality Assurance Project Plan. 991 Construction drawings and contract specifications will be 992 submitted to EPA and the State within three months from the entry of the Consent Decree. Comments provided by EPA and the State

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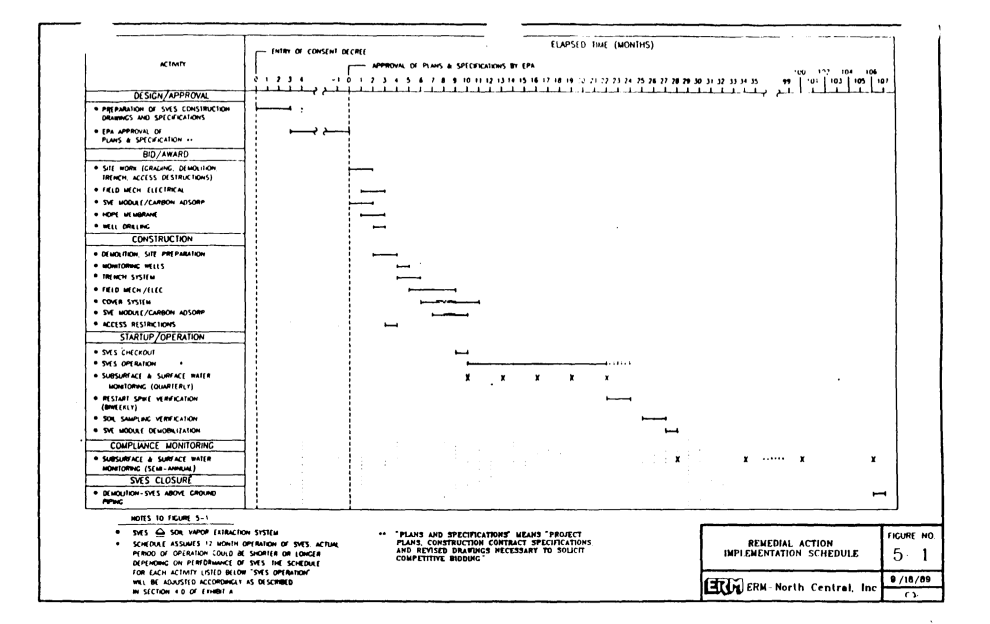
994 will be addressed by the Settling Defendants. 995 996 Figure 5-1 sets forth the Remedial Action Implementation Schedule 997 for implementing the remedy required under the Consent Decree. The following milestones have been established in Section XVII 998 999 (Stipulated Penalties) of the Consent Decree: 1000 1001 0 Submission of the project plans, construction 1002 contract specifications and revised drawings 1003 necessary to solicit competitive bidding 1004 within 3 months from the entry of the Decree. 1005 Completion of site preparation, including 1006 0 107 grading, removal of the tanks and buildings, **1008** repair or moving of the fence, 4 months after 1009 approval by EPA all of the above referenced 1010 documents. Completion of the site 1011 preparation shall mean that all hindrances. 1012 obstructions or obstacles to construction and 1013 security of the soil vapor extraction 1014 trenches, monitoring wells or cap have been 1015 removed. 1016 1017 Completion of installation of the on-site and 0 1018 off-site monitoring wells 5 months after approval by EPA of all of the above 1019 1020 referenced documents. 1021 1022 Startup of the soil vapor extraction system 0

10 months after approval by EPA of all of the

above referenced documents.

4 ∠5

1023



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1026	0	Completion of the installation of all	
1027		components of the RCRA-compliant (Subtitle C)	
1028		cover 11 months after approval by EPA of all	
1029		of the above referenced documents.	
1030			
1031	0	Submission of all documents necessary to	
1032		perform Additional Work that may be required	
1033		under Section VII of the Consent Decree 6	
1034		months after written notice has been provided	
1035		by EPA or Settling Defendants that Additional	
1036		Work needs to be implemented.	·
1037			
1038	0	Completion of installation of the subsurface	
1039		water interception trench on a schedule to be	
1040		determined by EPA after consultation with the	
1041		State.	

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YPPENDICES

APPENDIX A

ESTIMATE OF MASS OF ORGANICS IN THE SOILS TO BE REMOVED BY VAPOR EXTRACTION

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Location	Sampling depth (ft)	Assumed contamination depth (ft)	Total concentration (ug/kg)	Mass (lb)
TP-1 TP-2 TP-3 TP-4 TP-5 TP-5 TP-6 TP-6 TP-6 TP-6	1 - 1.5 1 - 1.5 1 - 1.5 1 - 2 2.5 - 3.5 1 - 2 2 - 3 1 - 2 2 - 3 4 - 5 1 - 2.5	2 2 2 2.5 4 2 1.5 2 1.5 1.5	1,972 28 108,800 99,730 4,416 24,287 291 12,468,000 22,690 2,416 267,000	0.271 0.004 14.978 17.162 1.216 3.343 0.030 1,716.410 2.343 0.249 45.946
TP-7 TP-8 TP-8 TP-9 TP-10 TP-10 TP-11 TP-11 TP-12 TP-12	2.5 - 4 1 - 2.5 2.5 - 4 1 - 3 3 - 5 1 - 3 3 - 5 1 - 3 3 - 5 1 - 3 3 - 5	2 2.5 2 3 2.5 3 2.5 3 2.5	280,090 3,687 433,600 14,604,000 130 958 432 130 67 35,030 3,609	38.559 0.634 59.692 3,015.694 0.022 0.198 0.074 0.027 0.012 7.234 0.621
SB-01 SB-02 SB-03 SB-04 SB-06 SB-08 SB-09 SB-01 SB-02 SB-04 SB-08 SB-09	2.5 - 4 2.5 - 4 2.5 - 4 2 - 3.5 2 - 3.5 2.5 - 4 2.5 - 4 5.5 - 7 5.5 - 7 5 - 6.5 7 - 8.5 5.7 - 7	3 3 2.5 2.5 3 3 2 2 2 2	3,303 12,900 70,070 175 222,010 3,012 61,490 27 34 51 188 8,069	0.682 2.664 14.469 0.030 38.204 0.622 12.698 0.004 0.005 0.007

TOTAL ORGANICS TO BE REMOVED BY VAPOR EXTRACTION, 1b 4,

^{*} The area contaminated is assumed to be a 25'x25' square around each sampling location. TP = test pit; SB = soil boring. Soil concentrations from ECC RI, Section 4.

APPENDIX B

CALCULATION OF RISK-BASED CLEANUP STANDARDS

HEARTH

APPENDIX B

CALCULATION OF RISK-BASED CLEANUP STANDARDS

The equations used to calculate risk-based concentrations are shown in Table B1. The ingestion rates and acceptable risks are listed in Table B2. The potency factors and references doses for compounds without any regulatory or background level are from a memorandum from the USEPA Toxics Integration Branch, OERR, Washington, D.C., dated December 19, 1988, with the Corrections to the July, 1988 Update of the Characterization Tables in the Superfund Public Health Evaluation Manual.

presents the calculation of risk-based acceptable subsurface water concentrations in the till for compounds without a regulatory limit (drinking water Maximum Contaminant Level, Maximum Contaminant Level Goal or lifetime health advisory or a stream criterion as listed in Table 1 of the Record of Decision Table B4 shows that the resulting concentrations for the site). of inorganic compounds at Unnamed Ditch should be below the Stream Criteria presented in Table 1 of the Record of Decision (ROD) for the site, dated September 25, 1987. obtained from discharge of the subsurface water in the till to Unnamed Ditch is 1:1800, as presented in Appendix C of the ECC Remedial Investigation. Note that most of the calculated concentrations in the ditch are below detection limits.

Tables B5 and B6 list the acceptable risk-based soil concentrations, based on soil and subsurface water ingestion, respectively. The calculation of acceptable soil concentrations based on subsurface water ingestion follows the procedures presented in Appendix C of the ECC RI. Only those organic compounds without regulatory limit (USEPA, Polychlorinated

MA.

TABLE 81 EQUATIONS USED TO CALCULATE RISK-BASED CONCENTRATIONS *

SOIL (concentrations in ug/kg):
Risk * Body Weight (kg) * 1000 (ug/mg) * 1000 (g/kg)
Ingestion rate (g/d) * Potency Factor (mg/kg/d)-1
or
Risk * Body Weight (kg) * Reference Dase (mg/kg/d) * 1000 (ug/mg) * 1000 (g/kg)
Ingestion rate (g/d)
SUBSURFACE WATER (concentrations in ug/l):
Risk * Body Weight (kg) * 1000 (ug/mg)
Ingestion rate (l/d) * Potency Factor (mg/kg/d)-1
ar
Risk * Body Weight (kg) * Reference Dose (mg/kg/d) * 1000 (ug/mg)
ingestion rate (l/d)

TABLE B2 INCESTION RATES AND ACCEPTABLE RISKS

INGESTION RATES . :

SOILS:

0.1 grams per day by a 70-kilogram person for 70 years

ar

0.2 grams per day by a 17-kilogram child for 5 years

SUBSURFACE WATER:

2 liters of water per day by a 70-kilogram person for 70 years

ACCEPTABLE RISKS:

COMPOUNDS WITH POTENCY FACTORS:

10

COMPOUNDS WITH REFERENCE DOSES:

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* From U.S. EPA, RCRA Facility Investigation Guidance, 1987, and U.S. EPA, Office of Solid Waste and Emergency Response, Hemorandum on Interim Final Guidance for Soil Ingestion Rates, January 27, 1989.

TABLE 83
ECC - ACCEPTABLE HEALTH-BASED SUBSURFACE WATER CONCENTRATIONS

Compound (1)	Potency Factor (2) (mg/kg/d)-1	Reference Dose (2) (mg/kg/d)	Acceptable Health-Based Subsurface water Concentration (3) (ug/l)
VOLATILE ORGANICS (VOCs):			
Acetone		0.1	3,500
1,1-Dichloroethane	0.091		0.38
Hethylene Chloride	0.0075		4.7
Methyl Isobutyl Ketone		0.05	1,750
Tetrachioroethene	0.051		0.69
1,1,2-Trichloroethane	0.057		0.61
BASE NEUTRAL/ACID ORGANICS:			
. Bis(2-ethylhexyl)phthalate	0.014		2.5
Di-n-Butyl Phthalate		0.1	3,500
Diethyl Phthalate		0.8	28,000
Isophorone	0.0041	•	8.5
Naphthalene		0.4	14,000
Phenol		0.04	1,400
PESTICIDES/PCBs:			
Aroctor-1232	7.7		0.0045
Aroctor-1260	7.7		0.0045
INORGANICS:			
Antimony		0.0004	14
Beryllium		0.005	175
Hanganese		0.2	7,000
Tin		0.6	21,000
Vanadius		0.007	245
Zinc		0.2	7,000

- (1) Only compounds without a regulatory limit (drinking water Maximum Contaminant Level [40 CFR 141], Maximum Contaminant Level Goal or lifetime health advisory) are shown.
- (2) From USEPA Toxics Integration Branch, OERR, Washington, D.C. December 1988 correction to the July 1988 Update of the Risk Characterization Tables in the Superfund Public Health Evaluation Manual.
- (3) Acceptable subsurface water concentrations calculated using an ingestion rate of 2 liters per day by a 70 kg adult for 70 years. Acceptable risk = 1E-06 for compounds with potency factor and 1 for compounds with reference dose.

TABLE B4 COMPARISON OF ACCEPTABLE STREAM CONCENTRATIONS WITH STREAM CONCENTRATIONS BASED ON NATURAL DISCHARGE OF SUBSURFACE WATER FROM THE TILL

Compounds (1)	Acceptable Stream Concentration (1) (ug/1)	Concentratic Unnamed Ditc to Discharge c Water at Accep Concentration (ug/l)
VOLATILE ORGANICS (VOCs): Chloroform 1,1-Dichloroethene Ethylbenzene Methylene Chloride Tetrachloroethene Toluene 1,1,1-Trichloroethane 1,1,2-Trichloroethane Trichloroethene	15.7 1.85 3,280 15.7 8.85 3,400 5,280 41.8 80.7	0.056 0.0039 1.9 0.0026 0.00038 5.8 0.11 0.00034 0.0028
BASE NEUTRAL/ACID ORGANICS: Bis(2-ethylhexyl)phthalate Di-n-Butyl Phthalate Diethyl Phthalate Naphthalene Phenol	50,000 154,000 52,100 620 570	0.0014 1.9 15.6 7.8 0.78
INORGANICS: Arsenic Chromium Lead Nickel Zinc Cyanide	0.0175 11 10 100 47 5.2	0.028 0.028 0.028 0.39 3.9 0.39

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From Table 1 of the Record of Decision (ROD) for the site, September 25, 1987. Only those compounds detected in ECC soil samples that are listed in this table are shown. Assuming a dilution of 1:1800 for natural discharge of till water at acceptable concentrations into Unnamed Ditch (from ECR Remedial Investigation, Appendix C). (2)

TABLE 85
ECC - ACCEPTABLE SOIL CONCENTRATIONS BASED ON SOIL INGESTION

Compounds (1)	Potency Factor (2) (mg/kg/d)-1	Reference Dase (2) (mg/kg/d)	Acceptable Soil Concentrations Based on Soil Ingestion (3) (ug/kg)	Range of Acceptable Soil Concentrations Based on Soil Ingestion (4) (ug/kg)
VOLATILE ORGANICS (VOCs):				
Acetone		0.1	8,500,000	8,500,000
Chiorobenzene		0.03	2,550,000	2,550,000
Chioroform	0.0061		114,754	11,475-11,475,400
1,1-Dichtoroethane	0.091		7,692	769-769,200
1,1-Dichloroethene	0.6		1,167	116.7-116,700
Ethylbenzene		0.1	8,500,000	8,500,000
Methylene Chloride	0.0075		93,333	9,333-9,333,300
Methyl Ethyl Ketone		0.05	4,250,000	~ ,250,000
Methyl Isobutyl Ketone		0.05	4,250,000	-,250,000 .
Tetrachloroethene	0.051		13,725	1,373-1,372,500
Toluene		0.3	25,500,000	25,500,000
1,1,1-Trichloroethane		0.09	7,650,000	7,650,000
1,1,2-Trichloroethane	0.057	•	12,281	1,228-1,228,100
Trichloroethene	0.011		63,636	6,364-6,363,600
Total Xylenes		2	170,000,000	. 170,000,000
BASE NEUTRAL/ACID ORGANICS:				
Bis(2-ethythexyt)phthatate	0.014		50,000	5,000-5,000,000
Di-n-Butyl Phthalate		0.1	8,500,000	8,500,000
Diethyl Phthalate		0.8	68,000,000	68,000,000
Isophorone	0.0041		170,732	17,073-17,073,200
Naphthalene		0.4	34,000,000	34,000,000
Phenoi		0.04	3,400,000	3,400,000

NOTES:

- (1) Only organic compounds without a regulatory limit in soils (USEPA, "Polychlorinated Biphenyls Spill Cleanup Policy Rule," 40 CFR Part 761) are shown.
- (2) From USEPA Toxics Integration Branch, DERR, Washington, D.C. December 19, 1988, "Corrections to the July 1988 Update of the Characterization Tables in the Superfund Public Health Evaluation Manual."
- (3) Intake for compounds with potency factor: 0.1 g of soil/d by 70 kg resident adults. Intake for compounds with reference dose: 0.2 g of soil/d by 17 kg resident children. Acceptable risks: 1E-06 for compounds with potency factor; 1 for compounds with reference dose.
- (4) Range shown is for risks of 10-4 to 10-7 for compounds with potency factor. The value shown for compounds without potency factor is for a risk of 1.

TABLE 86 (Page 1 of 2)
ECC - ACCEPTABLE SOIL CONCENTRATIONS BASED ON THEORETICAL SUBSURFACE WATER INGESTION AT THE SITE (10-6 RISK)

Compound (1)	\$olubility (2) (vg/l)	Log Kow (2)	Kd (3)	Acceptable Subsurface Water Concentration (4) (ug/l)	Acceptable Leachate Concentration (5) (ug/l)	Acceptable Soi Concentration Based on Water Ingestion (6) (ug/kg)
Acetone Chlorobenzene Chloroform 1,1-Dichloroethane 1,1-Dichloroethane 1,1-Dichloroethane Ethylbenzene Methylene Chloride Methyl Ethyl Ketone Methyl Isobutyl Katone Tetrachioroethane 1,1,1-Trichloroethane 1,1,2-Trichloroethane Trichloroethane Total Xylenes ASE NEUTRAL/ACID ORGANICS: Bis(2-ethylhexyl)phthalate Di-n-Butyl Phthalate Diethyl Phthalate Isophorone Maphthalene Phenol	1,000,000,000 466,000 8,200,000 5,500,000 2,250,000 152,000 20,000,000 268,000,000 17,000,000 200,000 535,000 4,400,000 4,500,000 1,100,000 198,000 1,300 13,000 12,000 30,000 93,000,000	-0.24 2.84 1.97 1.79 1.84 3.15 1.25 0.26 2.88 2.69 2.17 2.17 2.29 3.26 8.7 5.2 3.22	0.00071 0.858 0.116 0.076 0.086 1.75 0.022 0.00226 0.02604 0.941 0.607 0.183 0.183 0.242 2.26 621472 197 2.06 0.031 1.269 0.036	3,500 RB 60 MCLGP 100 MCL 0.38 RB 7 MCL 680 MCLGP 4.7 RB 170 LDWHA 1,750 RB 0.69 RB 2,000 MCLGP 200 MCL 0.61 RB 5 MCL 440 MCLGP 2.5 RB 3,500 RB 28,000RB 8.5 RB 14,000 RB 1,400 RB	686,275 11,765 19,608 74.5 1,373 133,333 922 33,333 343,137 135 392,157 39,216 120 980 86,275 490 686,275 5,490,196 1,667 2,745,098 274,510	490 10,093 2,269 5.7 118 233,540 20.3 75 8,935 127 238,167 7,193 21.9 237 194,672 304,643,220 134,871,303 11,298,207 51.7 3,483,209

TABLE 86 (Page 2 of 2)

ECC - ACCEPTABLE SOIL CONCENTRATIONS BASED ON THEORETICAL SUBSURFACE WATER INGESTION AT THE SITE (10-6 RISK)

- (1) Only organic compounds without a regulatory limit in soils (USEPA, "Polychlorinated Biphenyls Spill Cleanup Policy Rule," 40 CFR Part 761) are shown.
- (2) From ECC R1, Table 5-3, and Verschueren, 1983, "Mandbook of Environmental Data on Organic Chemicals".
- (3) From ECC RI, Table 5-3. Calculated as 10°log Kow * OC, where OC= organic carbon content = 0.00124. For isophorone and methyl isobutyl ketone, the Kd is obtained as Kd = Koc * OC, where Koc = organic carbon-water partition coefficient, obtained from log Koc = (-0.55 * log S) + 3.64 (Exhibit A-1 of *Superfund Public Health Evaluation Manual,* 1986).
- (4) RB = risk-based concentration, from Table B3; MCL = Maximum Contaminant Level, from 40 CFR 141; MCLGP = proposed MCL goal, from 40 CFR 141; LDWHA = lifetime drinking water health advisory, from "Superfund Public Health Evaluation Hanual," 1986.
- (5) Leachate discharge/subsurface water discharge = 0.0051 (Appendix C of the ECC RI; and reduction of the 7.8 in/yr recharge used in the RI under the current conditions (page 5-8) by 99 percent due to the cap).
- (6) Soil concentration (ug/kg) = Kd * Concentration in leachate (ug/l).

Biphenyls Spill Cleanup Police Rule, 40 CFR Part 761) in soils are listed in Tables B5 and B6. It is conservatively assumed that the volume of leachate from the soils will be reduced by 99 percent from the 7.8 in/yr used in the RI, by installing the RCRA-compliant (Subtitle C) cover over the site.

A range of acceptable soil concentrations based on water ingestion using the published ranges for organic carbon content of till soils and the SARA range of risk for Superfund site cleanups, is presented in Table B7. A list of organic carbon content in soil is shown in Table B8, with the respective reference. The concentrations shown in Table B6 were used to determine the Acceptable Soil Concentrations specified in Table 3-1, using a risk of 10⁻⁶ and a soil organic carbon content of 0.12%, as presented in the RI. This soil organic carbon content was deemed conservative when compared to the values shown in Table B8.

Table B9 lists the solubility and vapor pressure of the organic compounds detected in the soils above the limits shown in Tables B5 and B6. All compounds, except bis(2-ethylhexyl)phthalate and Aroclor-1260, are amenable to removal by soil vapor extraction.

Finally, Table B10 presents the complete list of references used for the calculation of the Acceptable Soil Concentrations specified in Table 3-1.

TABLE 87
ECC - ACCEPTABLE SOIL CONCENTRATIONS BASED ON THEORETICAL SUBSURFACE
- WATER INGESTION AT THE SITE (RANGE OF RISKS)

Acceptable Soil Concentration Based on Water Ingestion (3)

Compound (1)	Range of Kd (2)	Range for 10-4 risk	Range for 10-7 risk	
***********************	• • • • • • • • • • • • • • • • • • • •	***************************************	•••••••	
VOLATILE ORGANICS (VOCE):	•	•		
Acetone	0.000058-0.0044	40-3,019	40-3,019	(4)
Chiorobenzene	0.069-5.24	814-61,600	814-61,600	(5)
Chioroform	0.0093-0.71	182-13,900	182-13,900	(5)
1,1-Dichloroethane	0.0062-9_47	46-3,500	0.046-3.50	
1,1-Bichloroethene	0.0069-8.52	9.47-714	9.47-714	(5)
Ethylbenzene	0.14-10.7	18,800-1,431,000	18,800-1,431,000	(5)
Methylene Chloride	0.0018-0.14	166-12,900	0.166-12.9	
Methyl Ethyl Ketone	0.80018-0.014	6.07-461	6.07-461	(5)
Methyl Isobutyl Ketone	0 _0021-0.16	721-54,900	721-54,900	(4)
Tetrachioroethene	0.076-5.78	1,028-78,200	1.03-78.2-	
Taluene	0.049-3.72	19,200-1,460,000	19,200-1,460,000	(5)
1,1,1-Trichloroethane	0.015-1.14	588-44,700	588-44,700	(5)
1,1,2-Trichloroethane	0.015-1.14	179-13,600	0.179-13.6	
Trichloroethene	0.020-1.52	19.6-1,490	19.6-1,490	(5)
Total Xylenes	0.18-13.7	15,700-1,193,000	15,700-1,193,000	(5)
BASE NEUTRAL/ACID ORGANICS:				
#is(2-ethylhexyl)phthalate	50100-3810000	2,460,000,000-187,000,000,000	2,460,000-187,000,000	
Di-n-Butyl Phthalate	15.8-1200	10,800,000-824,000,000	10,800,000-824,000,000	(4)
Diethyl Phthalate	0.17-12.9	933,000-70,800,000	933,000-70,800,000	(4)
Isophorone	0.0025-0.19	417-31,700	0.417-31.7	
Naphthalene	0.1-7.6	275,000-20,900,000	275,000-20,900,000	(4)
Phenol	0.0029-0.22	796-60,400	796-60,400	(4)

NOTES:

- (1) Only organic compounds without a regulatory limit in soils (USEPA, "Polychlorinated Biphenyls Spill Cleanup Policy Rule," 40 CFR Part 761) are shown.
- (2) For a range of organic carbon content of 0.0001 to 0.0076 obtained from: U.S. Department of Agriculture, "Soil Classification - A Comprehensive System". Soil Conservation Service, 7th Approximation, 1960. Calculated as presented in Table 86.
- (3) Acceptable Soil Concentrations at the risk shown (for compounds with potency) for a range of organic carbon content of 0.0001 to 0.0076. Calculated as presented in Table 86.
- (4) Acceptable Soil Concentration range does not change because the compound does not have a potency factor.
- (5) Acceptable Soil Concentration range does not change because the value is based on regulatory limits (drinking water Maximum Contaminant Level, Maximum Contaminant Level Goal, or lifetime health advisory).

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TABLE 85 (Page 1 of 5) ORGANIC CARBON CONTENT OF SOILS - REFERENCES

Organic Carbon Content, %	Type of Soil (depth)	Geographic Area	Reference
0.125 (avg over 1.5 acre site)	Loamy sand (4 ft)	Etimanda, CA (arid region)	Elabd, H., and W.A. Jury. 1986. "Spatial variability of Pesticide Adsorption Parameters." Environmental Science and Technology, Vol. 20, No. 3, pp. 256-260.
0.2 (avg over 1.5 acre site)	Loamy sand (2 and 3 ft)	Ibid	1bi d
0.26 (avg over 1.5 acre site)	Loamy sand (1 ft)	Ibid	lbid
1.9	Silt loam	Corvallis, OR	Chiou, C.T., P.E. Porter, and D.W. Schmeddigh. 1983. "Partition Equilibria of Nonionic Organic Compounds between Soil Organic Matter and Water." Environmental Science and Technology, Vol. 17, No. 4, pp. 227-2314
0.15	Sand close to river	Switzerland	Schwarzenbach, R.P., and J. Westall. 1981. "Transport of Nonpolar Organic Compounds from Surface Water to Groundwater. Laboratory Sorption Studies." Environmental Science and Technology, Vol. 15, No. 11, pp. 1360-1367.
2.1	Air-dried soil	l owa	Wu, S., and P.M. Gschwend. 1986. "Sorption Kinetics of Hydrophobic Organic Compounds to Natural Sediments and Soils." Environmental Science and Technology, Vol. 20, No. 7, pp. 717-725.
0.11	Loess sample	Turin, lows	Karickhoff, S.W. 1984. **Organic Pollutant Sorption in Aquatic Systems.** Journal of Hydrautic Engineering, Vol. 110, No. 6, pp. 707-735.
1.3	Soil	Fern Clyffe State Park, IL	lbid
0.02	Aquifer water table zone 98 % sand	Borden, Canada	Abdul, A.S., T.L. Gibson, and D.N. Rai. 1986. "The Effect of Organic Carbon on the Adsorption of Fluorene by Aquifer Materials." Hazardous Waste and Mazardous Materials. Vol. 3, No. 4, pp. 429-440.
0.52	Aquifer water table zone 87 % sand	Flint, M1	1bid
1.8	Aquifer water table zone 91 % sand	Flint, M1	lbid

TABLE 88 (Page 2 of 5) ORGANIC CARBON CONTENT OF SOILS - REFERENCES

Organic Carbon Content, %	Type of Soil (depth)	Geographic Area	Reference
0.05	Fine-sand soil	Wilmington, DE	Stokman, S.K. 1987. "Estimates of Concentrations of Soluble Petroleum Hydrocarbons Migrating into Ground Water from Contaminated Soil Sources." Proceedings of the National Water Well Association/American Petroleum Institute Conference on Petroleum Hydrocarbons and Organic Chemicals in Ground Water - Prevention, Detection and Restoration. Houston, TX, pp. 541-558.
0.1	Shaly-silt soil	Philadelphia, PA	lbid
0.05	fine to coarse sand, 96% sand	Michigan	Chiang, C.Y., C.L. Klein, J.P. Salanitro, and H.L. Wisnieuski. 1986. "Data Analyses and Computer Modelling of the Benzene Plume in an Aquifer Beneath a Gas Plant." Proceedings of the National Water Well Association/American Petroleum Institute Conference ton Petroleum Hydrocarbons and Organic Chemicals in Ground Water - Prevention, Detection and Restoration. Houston, TX, pp. 157-176.
0.27	Lincoln fine sand (surface soil)	Little Sandy Creek near Ada, OK	Clark, G.L., A.T. Kan, and M.B. Tomson. 1986. "Kinetic Interaction of Neutral Trace Level Organic Compounds with Soil Organic Material." Proceedings of the National Water Well Association/American Petroleum Institute Conference on Petroleum Mydrocarbons and Organic Chemicals in Ground Water - Prevention, Detection and Restoration. Houston, TX, pp. 151-156.
0.74	Fine to medium grained sand (3 ft)	Indian River County, FL	Kemblowski, M.W., J.P. Salinatro, G.M. Deeley, and C.C. Stanley. 1987. "Fate and Transport of Residual Hydrocarbon in Groundwater - A Case Study." Proceedings of the National Water Well Association/ American Petroleum Institute Conference on Petroleum Hydrocarbons and Organic Chemicals in Ground Water - Prevention, Detection and Restoration. Houston, TX, pp. 207-231.
0.44	Fine to medium grained sand (7 ft)	Indian River County, FL	lbid
0.12	Fine to medium grained sand (13 ft)	Indian River County, FL	Ibid

TABLE 88 (Page 3 of 5) ORGANIC CARBON CONTENT OF SOILS - REFERENCES

Organic Carbon Content, %	Type of Soil (depth)	Geographic Area	Reference
0.36	Fine to medium grained sand (3 ft)	Indian River County, FL	lbid
0.15	Fine to medium grained sand (13 ft)	Indian River County, FL	lbid
1.08	Fine to medium grained sand (2 ft)	Indian River County, FL	Ibid
0.16	Fine to medium grained sand (11 ft)	Indian River County, FL	!bia
0.72	fine to medium grained sand (3 ft)	Indian River County, FL	i bi d
0.26	Fine to medium grained sand (10 ft)	Indian River County, FL	lbig ,
0.74	Glacial till (1-2 ft)	Sargent County, NO	MSoil Classification - A Comprehensive System.M 1960. U.S. Department of Agriculture, Soil Conservation Service, 7th Approximation.
0.33	Glacial till (2-3 ft)	Sargent County,	lbid
0.18	Glacial till (4.5-5 ft)	Sargent County, NO	Ibid
0.1	7fll (1-2 ft)	Strafford County, New Hampshire	lbíd
80.0	T((((2-3 ft)	Strafford County, New Hampshire	1bid
0.03	T{ (4-5 ft)	Strafford County, New Hampshire	1 b ı d
0.01	Till (5-7 ft)	Strafford County, New Hampshire	lbid

TABLE 88 (Page 4 of 5) ORGANIC CARBON CONTENT OF SOILS - REFERENCE

Organic Carbon Content, %	Type of Soil (depth)	Geographic Area
0.59	Calcareous,	
	glacial till	Greenbrier County,
	(1·2 ft)	West Virginia
0.27	Calcareous,	Connection
	glacial till	Greenbrier County, West Virginia
	(2-3 ft)	Seet Virginia
0.00		
0.08	Calcareous,	Greenbrier County,
	glacial till	West Virginia
	(4-5 ft)	••
0.38	Calassi	
	Calcareous,	Tomkins County,
	glacial till	New York
	(1-2 ft)	
0.16	Calcareous,	************************************
	glacial till	Tomkins County, New York
	(2-3 ft)	MEN TOLK
A 15		
0.17	Calcareous,	Tomkins County,
	glacial till	New York
	(4.5-6.5 ft)	
0.14	Calcareous,	•
	glacial till	Tomkins County,
	(6.5-7 ft)	New York
0.76	Glacial till	Haraca Calma
	(1-2 ft)	Waseca County, Minnesota
	•	WITH MESOCAL
0.3	Glacial till	Waseca County,
	(2-3 ft)	Minnesota
2.10		
0.19	Glacial till	Waseca County,
	(> 4 ft)	Mimesota
0.51	Glacial till	
		Sargent County,
	(1-2 ft)	MD
0.18	Glacial till	***
	(2-3 ft)	Sargent County,
		ND
0.16	Glacial till	Sacces of
	(3.5-5 ft)	Sargent County,
		ND

TABLE 88 (Page 5 of 5)
ORGANIC CARBON CONTENT OF SOILS - REFERENCES'

Organic Carbon Content, %	Type of Soil (depth)	Geographic Area	Reference
0.64	Firm, glacial till (1-2 ft)	Spink County, SD	lbid
0.36	Firm, glacial till (2-3 ft)	Spink County, SD	lbid
0.31	Firm, glacial till (4-5 ft)	Spink County, SD	Ibid
0.46	Glecial till (1-2 ft)	Renville County, NO	Ibid .
0.24	Glacial till (2-3 ft)	Renville County, ND	lbid .
0.13	Glacial till (4-5 ft)	Renville County, NO	Ibid
0.25	Glacial till (2-3 ft)	Adair County, Iowa	lbid
0.08	Glacial till (> 6 ft)	Adair County, lows	lbid
0.74	Calcareous, glacial till (1-2 ft)	Ward County, MD	1bid
0.2	Calcareous, glacial till (2-3 ft)	Ward County, MD	lbid!
0.19	Calcareous, glacial till (4-5 ft)	Ward County, NO	Ibid
0.35	Glacial till (1-2 ft)	Cayuga County, NY	1 bid
0.1	Glacial till (2-3 ft)	Cayuga County, NY	lbid
0.12	Glacial till (6-7 ft)	Cayuga County,	lbid

TABLE B9 CHEMICAL PROPERTIES OF ORGANIC COMPOUNDS DETECTED IN THE SOILS AT CONCENTRATIONS ABOVE THE ACCEPTABLE SOIL CONCENTRATIONS (1)

Compound	Solubility (ug/l)	Vapor Pressure (mm Hg)
VOLATILE ORGANICS (VOCs):		
Acetone	1,000,000,000	270
Chloroform	8,200,000	151
1,1-Dichloroethane	5,500,000	182
1,1-Dichloroethene	2,250,000	600
Ethylbenzene	152,000	7
Methylene Chloride	20,000,000	3 62 .
Methyl Ethyl Ketone	268,000,000	77.5
Methyl Isobutyl Ketone	17,000,000	6 .
Tetrachloroethene	200,000	17.8
Toluene	535,000	28.1
1,1,1-Trichloroethane	4,400,000	123
1,1,2-Trichloroethane	4,500,000	30
Trichloroethene	1,100,000	57.9
Total Xylenes	198,000	10
BASE NEUTRAL/ACID ORGANICS:	•	
Bis(2-ethylhexyl)phthalate	1,300	0.0000002
Isophorone	12,000	0.38
Phenol	93,000,000	0.341
PESTICIDES/PCBs:		
Aroclor-1260 (2)	2.7	0.0000405

- Acceptable Soil Concentrations are determined in accordance (1)
- with Footnotes 5, 6, and 7 of Table 3-1. Soil limit assumed for PCBs is 10,000 ug/kg (40 CFR Part (2) 761.125, "Polychlorinated Biphenyls Spill Cleanup Policy Rule").

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Pollutants," December 1979.

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USEPA, Toxics Integration Branch, OERR, Washington, D.C., December, 1988. "Memorandum with Corrections to the July, 1988 Update of the Risk Characterization Tables in the Superfund Public Health Evaluation Manual."

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APPENDIX C
ECC - VAPOR EXTRACTION MODEL

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APPENDIX C

ECC - VAPOR EXTRACTION MODEL

This program was written in FORTRAN by Michael C. Marley and George E. Hoag and reported in "Induced Soil Venting for Recovery/Restoration of Gasoline Hydrocarbons in the Vadose Zone," Proceedings, Petroleum Hydrocarbons and Organic Chemicals in Ground Water Conference, Houston, TX, 1984.

The program is based on the concentration of each component in the vapor phase in the soil, using the partial pressure exerted by each compound, as expressed by the following equation:

$$ZT = \frac{VP + X + V + MW}{R + T}$$

where:

ZT = concentration of the component in the vapor phase, mg/l

VP = vapor pressure of compound, mm Hg

X = mole fraction = moles of component/total moles of organics
in soil

V = volume of element, liters

MW = molecular weight of component

R = gas constant = 82.4 atm - cm³/gmoles^OK

 $T = temperature = 294.25^{\circ}K$

The program uses the finite difference method to calculate the change in number of moles of each component during a small time interval (i) and then recalculate over the next time interval (i+1), using the reduced number of moles resulting from subtracting the change in number of moles calculated for interval i from the number of moles present in the soil at the beginning of interval i.

The program runs for a finite length of time or until all the components are removed. The program was rewritten in BASIC and applied to the ECC site.

Table C-1 shows the chemical data used to run the model. The compounds to be evaluated are those shown in Table 3-2, which are amenable to removal by vapor extraction. The maximum detected soil concentrations were taken from Section 4 of the ECC RI, while the vapor pressure and molecular weight data are from USEPA, "Superfund Public Health Evaluation Manual," 1986.

As there was significant variation of compounds concentrations between soil samples at the site, a theoretical block size was chosen. This theoretical soil block is 10 ft x 10 ft x 2 ft deep and was assumed to contain all components of interest at their maximum detected concentrations (Table C-1). Furthermore, it was conservatively assumed that the air flow through the soil would only be 15% efficient in removing the organics. In effect, this represents a worst case estimate of the time required to remove the organics from the soils. The mass of this block was estimated as 10,200 kg.

TABLE C1
CHEMICAL DATA OF COMPOUNDS

	Molecular	Vapor Pressure (2)	Maximum Detected Soil Concentration (3		
Compound (1)	Weight (2)		(ug/kg)		
***************************************	*********	***********	************		
VOLATILE ORGANICS:					
Acetone	58.1	270	650,000		
Chloroform	119	151	2,900		
1,1-Dichloroethane	99	182	35,000		
1,1-Dichloroethene	97	600	380		
Ethylbenzene	106	7	1,500,000		
Methylene Chloride	85	362	310,000		
Methyl Ethyl Ketone	72.1	77.5	2,800,000		
Methyl Isobutyl Ketone	100	6	190,000		
Tetrachloroethene	166	17.8	650,00 0		
Toluene	92.1	28.1	2,000,000		
1,1,1-Trichloroethane	133	123	1,100,000		
1,1,2-Trichloroethane	133	30	550		
Trichloroethene	132	57.9	4,800,000		
BASE NEUTRAL/ACID ORGANICS:					
Phenol	94.1	0.341	570,000		
Isophorone	138	0.38	440,000		

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⁽¹⁾ Compounds shown are those amenable to soil vapor extraction.

⁽²⁾ From U.S. EPA, "Superfund Public Health Evaluation Manual," 1986.

⁽³⁾ From ECC RI, Harch 1986.

The air flow rate was estimated as a fraction of the total air flow rate to be used at the site (500 SCFM), based on the length of injection trench influencing the assumed soil block (10 ft) as a ratio of the total length of injection trenches (3,800 ft). This represents an air flow rate of 37.26 liters per minute.

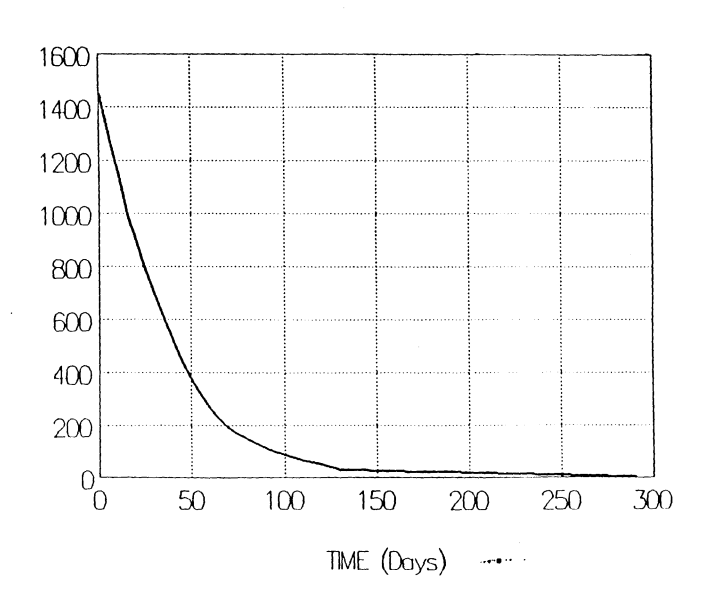
The results, summarized in Figure C1, show that essentially no VOCs will be present in the hypothetical soil element after 130 days of soil vapor extraction. To remove phenol and isophorone to the Acceptable Soil Concentrations in Table 3-1, operation of the vapor extraction system for a total of approximately 360 days is necessary.

Actual large-scale soil vapor extraction systems have been operated with excellent removals of compounds such as tetrachloroethene, trichloroethene, 1,3-dichloropropene, methyl ethyl ketone, methyl isobutyl ketone, toluene, and xylenes. Some published references are:

- o Lisiecki, J.B., and F.C. Payne. "Enhanced Volatilization: Possibilities, Practicalities, and Performance." Presented at the Engineering Foundation Conference, Mercersburg, PA, August 7-12, 1988.
- o Regalbuto, D.P., J.A. Barrera and J.B. Lisiecki. "In-Situ Removal of VOCs by Means of Enhanced Volatilization." Proceedings of the Conference on Petroleum Hydrocarbons and Organic Chemicals in Ground Water: Prevention, Detection, and Restoration, Houston, TX, November 9-11, 1988.

Figure 1

ECC VAPOR EXTRACTION MODEL RESULTS



o Johnson, J.J., and R.J. Sterrett. "Analysis of In-Situ Soil Air Stripping Data."

Proceedings of the 5th National Conference on Hazardous Wastes and Hazardous Materials, Las Vegas, Nevada, April 19-21, 1988.

A full-scale vapor extraction system (Lisiecki and Payne, 19 was able to remove tetrachloroethene from 5,600,000 ug/kg to ug/kg, as found by soil sample analysis, in 280 days. Therefo both theoretical models and actual results show that the requiremovals will be accomplished by vapor extraction.

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APPENDIX D

CALCULATION OF SOIL VAPOR CONCENTRATIONS

APPENDIX D

CALCULATION OF SOIL VAPOR CONCENTRATIONS

The methodology to determine the soil vapor concentrations in equilibrium with Acceptable Soil Concentrations in Table 3-1 is presented below.

The soil vapor concentration of a chemical in equilibrium with the concentration in the soil particles is a function of the soil to water partition coefficient and of the air to water partition coefficient [Lyman, W.J., W.F. Reehl and D.H. Rosenblatt, "Handbook of Chemical Property Estimation Methods," McGraw-Hill, Inc., 1982].

Since not all soil moisture will be evaporated during operation of the vapor extraction system (the soil's hygroscopic water will not be removed by the anticipated operating pressures), a relationship between soil vapor and soil moisture concentrations for the site's soils can be expressed as [Ibid] =

 $C_{SV} = H \cdot C_{SM}$

where:

 $C_{SV} = concentration of compound in soil vapor, mg/l$

H = Henry's Law Coefficient (nondimensional)

= <u>Vp . MW</u> S . R . T Vp = vapor pressure of compound, mm Hg

MW = molecular weight of the compound, g/gmole

s = solubility of the compound, g/cm³

R = gas law constant = $62,361 \text{ mm Hg} - \text{cm}^3/\text{gmole} - \text{OK}$

T = soil temperature = 283 OK

 C_{sm} = concentration of compound in soil moisture, mg/l

Similarly, the concentration in soil moisture in equilibrium with the concentration in soil particles can be calculated as [Ibid] =

where:

C_{sp} = concentration of compound in soil samples, mg/kg

Combining the two equations, a relationship between soil vapor and soil samples concentration is obtained [Silka, L.R., "Simulation of the Movement of Volatile Organic Vapor Through the Unsaturated Zone as it Pertains to Soil-Gas Surveys," Proceedings of the NWWA/API Conference on Petroleum Hydrocarbons and Organic Chemicals in Ground Water, 1986, p.204] =

$$C_{sv} = C_{sp} \cdot \frac{H}{K_d}$$

Table D1 presents the data and calculations of the soil vapiconcentration in equilibrium with the Acceptable Soil Concentrations in Table 3-1. None of the results shown in Table D1 is above the corresponding vapor saturation concentration, concentration in vapor in equilibrium with the pure compound The vapor saturation concentrations for the compounds in Table D1, assuming each compound is present by itself in the soil vapor (i.e., molar fraction is equal to 1), are shown in Table D2. The vapor saturation concentration is calculated as:

$$C_{sat} = \frac{Vp \cdot X \cdot MW}{R \cdot T} \times 10^6$$

where:

oour.

Csat = vapor saturation concentration, mg/l

X = molar fraction of compound in vapor, assumed to be 1

 10^6 = factor to convert g/cm^3 to mg/1

TABLE D1 (Page 1 of 2)

SOIL VAPOR CONCENTRATIONS IN EQUILIBRIUM
WITH ACCEPTABLE SOIL CONCENTRATIONS (1)

Compound (2)	Molecular Weight (3) (g/gmole)	Vapor Pressure (3) (mm Hg)	Solubility (3)	Henry's Law Constant (4) (dimensionless)	Soil-water Partition Coefficient (5)	Acceptable Soil Concentration (6)	Sai Canceni	l Vapor Tration (7)
VOLATILE ORGANICS (VOCs): Acetone			***********		(l/kg)	(ug/kg)	(mg/l)	ppmv
Chloroform 1,1-Dichloroethane 1,1-Dichloroethane 1,1-Dichloroethane Ethylbenzene Methylene Chloride Methyl Ethyl Ketone Methyl Isobutyl Ketone Tetrachloroethane 1,1,1-Trichloroethane 1,1,2-Trichloroethane Trichloroethane Total Xylenes SE NEUTRAL/ACID ORGANICS: Phenol	58.1 119 99 97 106 84.9 72.1 100 166 92.1 133 133 132 106	270 151 182 600 7 362 77.5 6 17.8 28.1 123 30 57.9	1,000,000,000 8,200,000 5,500,000 2,250,000 152,000 200,000 268,000,000 17,000,000 200,000 535,000 4,400,000 4,500,000 1,100,000 198,000	0.000889 0.124 0.186 1.47 0.277 0.0871 0.00118 0.00200 0.837 0.274 0.211 0.0502 0.394 0.303	0.00071 0.116 0.076 0.086 1.75 0.022 0.00226 0.026 0.941 0.607 0.183 0.183 0.242 2.26	490 2,300 5.7 120 234,000 20 -75 8,900 130 238,000 7,200 22 240 195,000	0.613 2.46 0.014 2.045 37 0.079 0.039 0.685 0.116 107 8.29 0.0060 0.39 26.2	254 496 3.39 515 9,316 22.4 13 233 16.8 36,556 2,819 1.09 71.5 4,794
	7.1	0.341	93,000,000	0.0000196	0.036	9,800	0.0053	1.36

.,,,,,,

TABLE D1 (Page 2 of 2) SOIL VAPOR CONCENTRATIONS IN EQUILIBRIUM WITH ACCEPTABLE SOIL CONCENTRATIONS (1)

NOTES:

- (1) Acceptable Soil Concentrations are determined in accordance with Footnotes 5 and 6 of Table 3-1.
- (2) Compounds above Acceptable Soil Concentrations in Table 3-1 to be removed by vapor extraction.
- (3) Data from U.S. EPA, "Superfund Public Nealth Evaluation Manual," 1986.
- (4) Calculated as:

(Vapor Pressure, mm Hg) * (Molecular Weight, g/gmole) * (1,000,000 ug/g) * (1,000 cm3/l)

Henry's Law Constant (nondimensional) =

(Solubility, ug/l) * (R, mm Hg-cm3/gmole-K) * (T, K)

where: R = gas law constant = 62,361 mm Hg-cm3/gmole-K; and T = soil temperature = 283 K.

- (5) From Appendix B, Table Bó.
- (6) From Table 3-1,
- (7) Calculated as:

(Concentration in soil, ug/kg) * (Henry's Law Constant, nondimensional)

Concentration in soil vapor (mg/l) *

(Partition coefficient, 1/kg) * (1000 ug/mg)

Concentration in soil vapor (ppmv) = (Concentration in soil vapor, mg/l) * (1000 l/m3) / (factor, mg/m3/ppmv)

The factors for conversion of mg/m3 to parts per million by volume (ppmv) were obtained from Vershueren, K., "Mandbook of Environmental Environmental Data on Organic Chemicals," 2nd Edition, 1983.

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TABLE D2
CALCULATION OF VAPOR SATURATION CONCENTRATIONS

Compound (1)	Vapor Pressure (2) (mm Hg)	Molecular Weight (2) (g/gmole)	Vapor Saturation Concentration (3) (mg/l)
		•••••	***************************************
VOLATILE ORGANICS (VOCs):			
Acetone	270	58.1	888.9
Chloroform	151	119	1018.2
1,1-Dichloroethane	182	99	1021.0
1,1-Dichloroethene	600	97	3297.8
Ethylbenzene	7	106	42.0
Methylene Chloride	362	84.9	1741.5
Methyl Ethyl Ketone	<i>7</i> 7.5	72.1	316.6
Methyl Isobutyl Ketone	6	100	34.0
Tetrachloroethene	17.8	166	167.4
Toluene	28.1	92.1	146.6
1,1,1-Trichloroethane	123	133	927.0
1,1,2-Trichloroethane	30	133	226.1
Trichloroethene	57.9	132	433.1
Total Xylenes	10	106	60.1
BASE NEUTRAL/ACID ORGANICS:			
Phenal	0.341	94.1	1.8

Where: Csat = vapor saturation concentration, mg/l; X = motar fraction of compound in vapor, assumed to be 1; 1E+06 = factor to convert g/cm3 to mg/l; MW = molecular weight of the compound, g/gmole; R = gas law constant, 62,361 mm Hg-cm3/gmole-K; and T = soil temperature, 283 K.

⁽¹⁾ Compounds above Acceptable Soil Concentrations in Table 3-1 to be removed by vapor extraction.

⁽²⁾ Data from U.S. EPA, "Superfund Public Health Evaluation Manual," 1986.

⁽³⁾ Calculated as: